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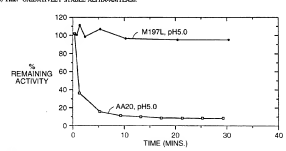
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(54) Title: OXIDATIVELY STABLE ALPHA-AMYLASE



# (57) Abstract

Novel alpha-amylase mutants derived from the DNA sequences of naturally occurring or recombinant alpha-amylases are disclosed. The mutant alpha-amylases in general, are obtained by nivro modifications of a preusors DNA sequence encoding the naturally occurring or recombinant alpha-amylase to generate the substitution (replacement) or deletion of one or more oxidizable amino acid residues in the amino acid sequence of a precursor alpha-amylase. Such mutant alpha-amylases have altered oxidative stability and relatered pH performance profiles and/or altered thermal stability as compared to the precursor. Also disclosed are detergent and starch liquefaction compositions comprising the mutant amylases, as well as methods of using the mutant amylases, as

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## OXIDATIVELY STABLE ALPHA-AMYLASE

## Related Applications

This application is a continuation-in-part of USSN 08/016,395 filed February 11, 1993.

## Field of the Invention

The present invention relates to novel alpha-amylase mutants having an amino acid sequence not found in nature, such mutants having an amino acid sequence wherein one or more amino acid residue(s) of a precursor alpha-amylase, specifically any oxidizable amino acid, have been substituted with a different amino acid. The mutant enzymes of the present invention exhibit altered stability/activity profiles including but not limited to altered oxidative stability, altered pH performance profile, altered specific activity and/or altered thermostability.

## Background of the Invention

Alpha-amylases (alpha-1,4-glucan-4-glucanohydrolase, EC3.2.1.1) hydrolyze internal alpha-1,4-glucosidic linkages in starch largely at random, to produce smaller molecular weight malto-dextrins. Alpha-amylases are of considerable commercial value, being used in the initial stages (liquefaction) of starch processing; in alcohol production; as cleaning agents in detergent matrices; and in the textile industry for starch desizing. Alpha-amylases are produced by a wide variety of microorganisms including Bacillus and Aspergillus, with most commercial amylases being produced from bacterial sources such as B. licheniformis, B. amyloliquefaciens, B. subtilis, or B. stearothermophilus. In

recent years the preferred enzymes in commercial use have been those from B.

licheniformis because of their heat stability and performance, at least at neutral and mildly alkaline pH's.

Previously there have been studies using recombinant DNA techniques to explore which residues are important for the catalytic activity of amylases and/or to explore the effect of modifying certain amino acids within the active site of various amylases (Vihinen, M. et al. (1990) J. Bichem. 107:267-272; Holm, L. et al. (1990) Protein Engineering 3:181-191; Takase, K. et al. (1992) Biochemica et Biophysica Acta, 1120:281-288; Matsui, I. et al. (1992) Febs Letters Vol. 310, No. 3, pp. 216-218); which residues are important for thermal stability (Suzuki, Y. et al. (1989) J. Biol. Chem. 264:18933-18938); and one group has used such methods to introduce mutations at various histidine residues in a *B. licheniformis* amylase, the rationale for making substitutions at histidine residues was that *B. licheniformis* amylase (known to be thermostable) when compared to other similar *Bacillus* amylases, has an excess of histidines and, therefore, it was suggested that replacing a histidine could affect the thermostability of the enzyme (Declerck, N. et al. (1990) J. Biol. Chem. 265:15481-15488; FR 2 665 178-A1; Joyet, P. et al. (1992) Bio/Technology 10:1579-1583).

It has been found that alpha-amylase is inactivated by hydrogen peroxide and other oxidants at pH's between 4 and 10.5 as described in the examples herein.

Commercially, alpha-amylase enzymes can be used under dramatically different conditions such as both high and low pH conditions, depending on the commercial application. For example, alpha-amylases may be used in the liquefaction of starch, a process preferably performed at a low pH (pH < 5.5). On the other hand, amylases may be used in commercial dish care or laundry detergents, which often contain

oxidants such as bleach or peracids, and which are used in much more alkaline conditions.

In order to alter the stability or activity profile of amylase enzymes under varying conditions, it has been found that selective replacement, substitution or deletion of oxidizable amino acids, such as a methionine, tryptophan, tyrosine, histidine or cysteine, results in an altered profile of the variant enzyme as compared to its precursor. Because currently commercially available amylases are not acceptable (stable) under various conditions, there is a need for an amylase having an altered stability and/or activity profile. This altered stability (oxidative, thermal or pH performance profile) can be achieved while maintaining adequate enzymatic activity, as compared to the wild-type or precursor enzyme. The characteristic affected by introducing such mutations may be a change in oxidative stability while maintaining thermal stability or vice versa. Additionally, the substitution of different amino acids for an exidizable amino acids in the alpha-amylase precursor sequence or the deletion of one or more oxidizable amino acid(s) may result in altered enzymatic activity at a pH other than that which is considered optimal for the precursor alpha-amylase. In other words, the mutant enzymes of the present invention may also have altered pH performance profiles, which may be due to the enhanced oxidative stability of the enzyme.

#### Summary of the Invention

The present invention relates to novel alpha-amylase mutants that are the expression product of a mutated DNA sequence encoding an alpha-amylase, the mutated DNA sequence being derived from a precursor alpha-amylase by the deletion or substitution (replacement) of one or more oxidizable amino acid. In one preferred embodiment of

the present invention the mutant result from substituting a different amino acid for one or more methionine residue(s) in the precursor alpha-amylase. In another embodiment of the present invention the mutants comprise a substitution of one or more tryptophan residue alone or in combination with the substitution of one or more methionine residue in the precursor alpha-amylase. Such mutant alpha-amylases, in general, are obtained by *in vitro* modification of a precursor DNA sequence encoding a naturally occurring or recombinant alpha-amylase to encode the substitution or deletion of one or more amino acid residues in a precursor amino acid sequence.

Preferably the substitution or deletion of one or more amino acid in the amino acid sequence is due to the replacement or deletion of one or more methionine, tryptophan, cysteine, histidine or tyrosine residues in such sequence, most preferably the residue which is changed is a methionine residue. The oxidizable amino acid residues may be replaced by any of the other 20 naturally occurring amino acids. If the desired effect is to alter the oxidative stability of the precursor, the amino acid residue may be substituted with a non-oxidizable amino acid (such as alanine, arginine, asparagine, aspartic acid, glutamic acid, glutamine, glycine, isoleucine, leucine, lysine, phenylalanine, proline, serine, threonine, or valine) or another oxidizable amino acid (such as cysteine, methionine, tryptophan, tyrosine or histidine, listed in order of most easily oxidizable to less readily oxidizable). Likewise, if the desired effect is to alter thermostability, any of the other 20 naturally occurring amino acids may be substituted (i.e., cysteine may be substituted for methionine).

Preferred mutants comprise the substitution of a methionine residue equivalent to any of the methionine residues found in *B. licheniformis* alpha-amylase (+8, +15, +197, +256, +304, +366 and +438). Most preferably the methionine to be replaced is a

methionine at a position equivalent to position +197 or +15 in *B. licheniformis* alphaamylase. Preferred substitute amino acids to replace the methionine at position +197 are alanine (A), isoleucine (I), threonine (T) or cysteine (C). The preferred substitute amino acids at position +15 are leucine (L), threonine (T), asparagine (N), aspartate (D), serine (S), valine (V) and isoleucine (I), although other substitute amino acids not specified above may be useful. Two specifically preferred mutants of the present invention are M197T and M15L.

Another embodiment of this invention relates to mutants comprising the substitution of a tryptophan residue equivalent to any of the tryptophan residues found in *B. licheniformis* alpha-amylase (see Fig. 2). Preferably the tryptophan to be replaced is at a position equivalent to +138 in *B. licheniformis* alpha-amylase. A mutation (substitution) at a tryptophan residue may be made alone or in combination with mutations at other oxidizable amino acid residues. Specifically, it may be advantageous to modify by substitution at least one tryptophan in combination with at least one methionine (for example, the double mutant +138/+197).

The alpha-amylase mutants of the present invention, in general, exhibit altered oxidative stability in the presence of hydrogen peroxide and other oxidants such as bleach or peracids, or, more specific, milder oxidants such as chloramine-T. Mutant enzymes having enhanced oxidative stability will be useful in extending the shelf life and bleach, perborate, percarbonate or peracid compatibility of amylases used in cleaning products. Similarly, reduced oxidative stability may be useful in industrial processes that require the rapid and efficient quenching of enzymatic activity. The mutant enzymes of the present invention may also demonstrate a broadened pH performance profile whereby mutants such as M15L show stability for low pH starch

liquefaction and mutants such as M197T show stability at high pH cleaning product conditions. The mutants of the present invention may also have altered thermal stability whereby the mutant may have enhanced stability at either high or low temperatures. It is understood that any change (increase or decrease) in the mutant's enzymatic characteristic(s), as compared to its precursor, may be beneficial depending on the desired end use of the mutant alpha-amylase.

In addition to starch processing and cleaning applications, variant amylases of the present invention may be used in any application in which known amylases are used, for example, variant amylases can be used in textile processing, food processing, etc. Specifically, it is contemplated that a variant enzyme such as M197C, which is easily inactivated by oxidation, would be useful in a process where it is desirable to completely remove amylase activity at the end of the process, for example, in frozen food processing applications.

The preferred alpha-amylase mutants of the present invention are derived from a Bacillus strain such as B. licheniformis, B. amyloliquefaciens, and B. stearothermophilus, and most preferably from Bacillus licheniformis.

In another aspect of the present invention there is provided a novel form of the alphaamylase normally produced by *B. licheniformis*. This novel form, designated as the A4 form, has an additional four alanine residues at the N-terminus of the secreted amylase. (Fig. 4b.) Derivatives or mutants of the A4 form of alpha-amylase are encompassed within the present invention. By derivatives or mutants of the A4 form, it is meant that the present invention comprises the A4 form alpha-amylase containing one or more additional mutations such as, for example, mutation (substitution, replacement or

deletion) of one or more oxidizable amino acid(s).

In a composition embodiment of the present invention there are provided detergent compositions, liquid, gel or granular, comprising the alpha-amylase mutants described herein. Particularly preferred are detergent compositions comprising a +197 position mutant either alone or in combination with other enzymes such as endoglycosidases, cellulases, proteases, lipases or other amylase enzymes. Additionally, it is contemplated that the compositions of the present invention may include an alpha-amylase mutant having more than one site-specific mutation.

In yet another composition embodiment of the present invention there are provided compositions useful in starch processing and particularly starch liquefaction. The starch liquefaction compositions of the present invention preferably comprise an alphaamylase mutant having a substitution or deletion at position M15. Additionally, it is contemplated that such compositions may comprise additional components as known to those skilled in the art, including, for example, antioxidents, calcium, ions, etc.

In a process aspect of the present invention there are provided methods for liquefying starch, and particularly granular starch slurries, from either a wet or dry milled process. Generally, in the first step of the starch degradation process, the starch slurry is gelatinized by heating at a relatively high temperature (up to about 110°C). After the starch slurry is gelatinized it is liquefied and dextrinized using an alpha-amylase. The conditions for such liquefaction are described in commonly assigned US patent applications 07/785,624 and 07/785,623 and US Patent 5,180,669, the disclosure of which are incorporated herein by reference. The present method for liquefying starch comprises adding to a starch slurry an effective amount of an alpha-amylase of the

present invention, alone or in combination with additional excipients such as an antioxidant, and reacting the slurry for an appropriate time and temperature to liquefy the starch.

A further aspect of the present invention comprises the DNA encoding the mutant alpha-amylases of the present invention (including A4 form and mutants thereof) and expression vectors encoding the DNA as well as host cells transformed with such expression vectors.

## Brief Description of the Drawings

Fig. 1 shows the DNA sequence of the gene for alpha-amylase from *B. lichenitormis* (NCIB8061), Seq ID No 31, and deduced translation product as described in Gray, G. et al. (1986) J. Bacter. 166:635-643.

Fig. 2 shows the amino acid sequence of the mature alpha-amylase enzyme from *B. licheniformis* (NCIB8061), Seq ID No 32.

Fig. 3 shows an alignment of primary structures of *Bacillus* alpha-amylases. The *B. licheniformis* amylase (Am-Lich), Seq ID No 33, is described by Gray, G. et al. (1986)

J. Bact. 166:635-643; the *B. amyloliquefaciens* amylase (Am-Amylo), Seq ID No 34, is described by Takkinen, K. et al. (1983) J. Biol. Chem. 258:1007-1013; and the *B. stearothermophilus* (Am-Stearo), Seq ID No 35, is described by Ihara, H. et al. (1985)

J. Biochem. 98:95-103.

Fig. 4a shows the amino acid sequence of the mature alpha-amylase variant M197T, Sea ID No 36.

Fig. 4b shows the amino acid sequence of the A4 form of alpha-amylase from *B. licheniformis* NCIB8061, Seq ID No 37. Numbering is from the N-terminus, starting with the four additional alanines.

Fig. 5 shows plasmid pA4BL wherein BLAA refers to *B. licheniformis* alpha-amylase gene, Pstl to Sstl; Amp<sup>8</sup> refers to the ampicillin-resistant gene from pBR322; and CAT refers to the Chloramphenicol-resistant gene from pC194.

Fig. 6 shows the signal sequence-mature protein junctions for *B. licheniformis* (Seq ID No 38), *B. subtilis* (Seq ID No 39), *B. licheniformis* in pA4BL (Seq ID No 40) and *B. licheniformis* in pBLapr (Sea ID No 41).

Fig. 7a shows inactivation of certain alpha-amylases (Spezyme® AA20 and M197L (A4 form) with 0.88M H<sub>2</sub>O<sub>2</sub> at pH 5.0, 25 °C.

Fig. 7b shows inactivation of certain alpha-amylases (Spezyme® AA20, M197T) with 0.88M H<sub>2</sub>O<sub>2</sub> at pH 10.0, 25°C.

Fig. 7c shows inactivation of certain alpha-amylases (Spezyme® AA20, M15L) with 0.88M H<sub>2</sub>O<sub>2</sub> at pH 5.0, 25°C.

Fig. 8 shows a schematic for the production of M197X cassette mutants.

Fig. 9 shows expression of M197X variants.

Fig. 10 shows thermal stability of M197X variants at pH 5.0, 5mM CaCl<sub>2</sub> at 95°C for 5

mins.

Figs. 11a and 11b show inactivation of certain amylases in automatic dish care detergents. Fig. 11a shows the stability of certain amylases in Cascade<sup>™</sup> (a commercially available dish care product) at 65°C in the presence or absence of starch. Fig. 11b shows the stability of certain amylases in Sunlight<sup>™</sup> (a commercially available dish care product) at 65°C in the presence or absence of starch.

Fig. 12 shows a schematic for the production of M15X cassette mutants.

Fig. 13 shows expression of M15X variants.

Fig. 14 shows specific activity of M15X variants on soluble starch.

Fig. 15 shows heat stability of M15X variants at 90°C, pH 5.0, 5mM CaCl<sub>2</sub>, 5 mins.

Fig. 16 shows specific activity on starch and soluble substrate, and performance in jet liquefaction at pH 5.5, of M15 variants as a function of percent activity of *B. licheniformis* wild-type.

Fig. 17 shows the inactivation of *B. licheniformis* alpha-amylase (AA20 at 0.65 mg/ml) with chloramine-T at pH 8.0 as compared to variants M197A (1.7 mg/ml) and M197L (1.7 mg/ml).

Fig. 18 shows the inactivation of *B. licheniformis* alpha-amylase (AA20 at 0.22 mg/ml) with chloramine-T at pH 4.0 as compared to variants M197A (4.3 mg/ml) and M197L

(0.53 ma/ml).

Fig. 19 shows the reaction of *B. licheniformis* alpha-amylase (AA20 at 0.75 mg/ml) with chloramine-T at pH 5.0 as compared to double variants M197T/W138F (0.64 mg/ml) and M197T/W138Y (0.60 mg/ml).

## Detailed Description of the Invention

It is believed that amylases used in starch liquefaction may be subject to some form of inactivation due to some activity present in the starch slurry (see commonly owned US applications 07/785,624 and 07/785,623 and US Patent 5,180,669, issued January 19, 1993, incorporated herein by reference). Furthermore, use of an amylase in the presence of oxidants, such as in bleach or peracid containing detergents, may result in partial or complete inactivation of the amylase. Therefore, the present invention focuses on altering the oxidative sensitivity of amylases. The mutant enzymes of the present invention may also have an altered pH profile and/or altered thermal stability which may be due to the enhanced oxidative stability of the enzyme at low or high pH's.

Alpha-amylase as used herein includes naturally occurring amylases as well as recombinant amylases. Preferred amylases in the present invention are alpha-amylases derived from *B. licheniformis* or *B. stearothermophilus*, including the A4 form of alpha-amylase derived from *B. licheniformis* as described herein, as well as fungal alpha-amylases as those derived from *Aspergillus* (i.e. as *A. oryzae* and *A. niger*).

Recombinant alpha-amylases refers to an alpha-amylase in which the DNA sequence encoding the naturally occurring alpha-amylase is modified to produce a mutant DNA

sequence which encodes the substitution, insertion or deletion of one or more amino acids in the alpha-amylase sequence. Suitable modification methods are disclosed herein, and also in commonly owned US Patents 4,760,025 and 5,185,258, the disclosure of which are incorporated herein by reference.

Homologies have been found between almost all endo-amylases sequenced to date, ranging from plants, mammals, and bacteria (Nakajima, R.T. et al. (1986) Appl. Microbiol, Biotechnol. 23:355-360; Rogers, J.C. (1985) Biochem. Biophys. Res. Commun. 128:470-476). There are four areas of particularly high homology in certain Bacillus amylases, as shown in Fig. 3, wherein the underlined sections designate the areas of high homology. Further, sequence alignments have been used to map the relationship between Bacillus endo-amylases (Feng. D.F. and Doolittle, R.F. (1987) J. Molec. Evol. 35:351-360). The relative sequence homology between B. stearothermophilus and B. licheniformis amylase is about 66%, as determined by Holm, L. et al. (1990) Protein Engineering 3 (3) pp. 181-191. The sequence homology between B. licheniformis and B. amyloliquefaciens amylases is about 81%, as per Holm, L. et al., supra. While sequence homology is important, it is generally recognized that structural homology is also important in comparing amylases or other enzymes. For example, structural homology between fungal amylases and bacterial (Bacillus) amylase have been suggested and, therefore, fungal amylases are encompassed within the present invention.

An alpha-amylase mutant has an amino acid sequence which is derived from the amino acid sequence of a precursor alpha-amylase. The precursor alpha-amylases include naturally occurring alpha-amylases and recombinant alpha-amylases (as defined). The amino acid sequence of the alpha-amylase mutant is derived from the precursor alpha-amylase mutant is derived from the precursor alpha-amylase.

amylase amino acid sequence by the substitution, deletion or insertion of one or more amino acids of the precursor amino acid sequence. Such modification is of the precursor DNA sequence which encodes the amino acid sequence of the precursor alpha-amylase rather than manipulation of the precursor alpha-amylase enzyme per se. Suitable methods for such manipulation of the precursor DNA sequence include methods disclosed herein and in commonly owned US patent 4,760,025 and 5,185,258.

Specific residues corresponding to positions M197, M15 and W138 of *Bacillus*licheniformis alpha-amylase are identified herein for substitution or deletion, as are all methionine, histidine, tryptophan, cysteine and tyrosine positions. The amino acid position number (i.e., +197) refers to the number assigned to the mature *Bacillus*licheniformis alpha-amylase sequence presented in Fig. 2. The invention, however, is not limited to the mutation of this particular mature alpha-amylase (*B. licheniformis*) but extends to precursor alpha-amylases containing amino acid residues at positions which are equivalent to the particular identified residue in *B. licheniformis* alpha-amylase. A residue (amino acid) of a precursor alpha-amylase is equivalent to a residue of *B. licheniformis* alpha-amylase if it is either homologous (i.e., corresponding in position in either primary or tertiary structure) or analogous to a specific residue or portion of that residue in *B. licheniformis* alpha-amylase (i.e., having the same or similar functional capacity to combine, react, or interact chemically or structurally).

In order to establish homology to primary structure, the amino acid sequence of a precursor alpha-amylase is directly compared to the *B. licheniformis* alpha-amylase primary sequence and particularly to a set of residues known to be invariant to all alpha-amylases for which sequence is known, as seen in Fig. 3. It is possible also to

determine equivalent residues by tertiary structure: crystal structures have been reported for porcine pancreatic alpha-amylase (Buisson, G. et al. (1987) EMBO J.6:3909-3916); Taka-amylase A from Aspergillus oryzae (Matsuura, Y. et al. (1984) J. Biochem. (Tokyo) 95:697-702); and an acid alpha-amylase from A. niger (Boel, E. et al. (1990) Biochemistry 29:6244-6249), with the former two structures being similar. There are no published structures for Bacillus alpha-amylases, although there are predicted to be common super-secondary structures between glucanases (MacGregor, E.A. & Svensson, B. (1989) Biochem. J. 259:145-152) and a structure for the B. stearothermophilus enzyme has been modeled on that of Taka-amylase A (Holm, L. et al. (1990) Protein Engineering 3:181-191). The four highly conserved regions shown in Fig. 3 contain many residues thought to be part of the active-site (Matsuura, Y. et al. (1984) J. Biochem. (Tokyo) 95:697-702; Buisson, G. et al. (1987) EMBO J. 6:3909-3916; Vihinen, M. et al. (1990) J. Biochem. 107:267-272) including, in the licheniformis numbering, His105; Arg229; Asp231; His235; Glu261 and Asp328.

Expression vector as used herein refers to a DNA construct containing a DNA sequence which is operably linked to a suitable control sequence capable of effecting the expression of said DNA in a suitable host. Such control sequences may include a promoter to effect transcription, an optional operator sequence to control such transcription, a sequence encoding suitable mRNA ribosome-binding sites, and sequences which control termination of transcription and translation. A preferred promoter is the *B. subtilis aprE* promoter. The vector may be a plasmid, a phage particle, or simply a potential genomic insert. Once transformed into a suitable host, the vector may replicate and function independently of the host genome, or may, in some instances, integrate into the genome itself. In the present specification, plasmid and vector are sometimes used interchangeably as the plasmid is the most commonly

used form of vector at present. However, the invention is intended to include such other forms of expression vectors which serve equivalent functions and which are, or become, known in the art.

Host strains (or cells) useful in the present invention generally are procaryotic or eucaryotic hosts and include any transformable microorganism in which the expression of alpha-amylase can be achieved. Specifically, host strains of the same species or genus from which the alpha-amylase is derived are suitable, such as a *Bacillus* strain. Preferably an alpha-amylase negative *Bacillus* strain (genes deleted) and/or an alpha-amylase and protease deleted *Bacillus* strain such as *Bacillus* subtilis strain BG2473 (Δ*amyE*, Δ*apr*, Δ*npr*) is used. Host cells are transformed or transfected with vectors constructed using recombinant DNA techniques. Such transformed host cells are capable of either replicating vectors encoding the alpha-amylase and its variants (mutants) or expressing the desired alpha-amylase.

Preferably the mutants of the present invention are secreted into the culture medium during fermentation. Any suitable signal sequence, such as the *aprE* signal peptide, can be used to achieve secretion.

Many of the alpha-amylase mutants of the present invention are useful in formulating various detergent compositions, particularly certain dish care cleaning compositions, especially those cleaning compositions containing known oxidants. Alpha-amylase mutants of the invention can be formulated into known powdered, liquid or gel detergents having pH between 6.5 to 12.0. Suitable granular composition may be made as described in commonly owned US patent applications 07/429,881, 07/533,721 and 07/957,973, all of which are incorporated herein by reference. These

detergent cleaning compositions can also contain other enzymes, such as known proteases, lipases, cellulases, endoglycosidases or other amylases, as well as builders, stabilizers or other excipients known to those skilled in the art. These enzymes can be present as co-granules or as blended mixes or in any other manner known to those skilled in the art. Furthermore, it is contemplated by the present invention that multiple mutants may be useful in cleaning or other applications. For example, a mutant enzyme having changes at both +15 and +197 may exhibit enhanced performance useful in a cleaning product or a multiple mutant comprising changes at +197 and +138 may have improved performance.

As described previously, alpha-amylase mutants of the present invention may also be useful in the liquefaction of starch. Starch liquefaction, particularly granular starch slurry liquefaction, is typically carried out at near neutral pH's and high temperatures. As described in commonly owned US applications 07/788,624 and 07/785,623 and US Patent 5,180,669, it appears that an oxidizing agent or inactivating agent of some sort is also present in typical liquefaction processes, which may affect the enzyme activity; thus, in these related patent applications an antioxidant is added to the process to protect the enzyme.

Based on the conditions of a preferred liquefaction process, as described in commonly owned US applications 07/788,624 and 07/785,623 and US Patent 5,180,669, namely low pH, high temperature and potential oxidation conditions, preferred mutants of the present invention for use in liquefaction processes comprise mutants exhibiting altered pH performance profiles (i.e., low pH profile, pH <6 and preferably pH <5.5), and/or altered thermal stability (i.e., high temperature, about 90°-110°C), and/or altered oxidative stability (i.e., enhanced oxidative stability).

Thus, an improved method for liquefying starch is taught by the present invention, the method comprising liquefying a granular starch slurry from either a wet or dry milling process at a pH from about 4 to 6 by adding an effective amount of an alpha-amylase mutant of the present invention to the starch slurry; optionally adding an effective amount of an antioxidant or other excipient to the slurry; and reacting the slurry for an appropriate time and temperature to liquefy the starch.

The following is presented by way of example and is not to be construed as a limitation to the scope of the claims. Abbreviations used herein, particularly three letter or one letter notations for amino acids are described in Dale, J.W., Molecular Genetics of Bacteria. John Wiley & Sons. (1989) Appendix B.

## Experimental

BclI

## Example 1

Substitutions for the Methionine Residues in B. licheniformis Alpha-Amylase

The alpha-amylase gene (Fig. 1) was cloned from B. licheniformis NCIB8061 obtained
from the National Collection of Industrial Bacteria, Aberdeen, Scotland (Gray, G. et al.
(1986) J. Bacteriology 166:635-643). The 1.72kb Psti-Sstl fragment, encoding the
last three residues of the signal sequence; the entire mature protein and the terminator
region was subcloned into M13MP18. A synthetic terminator was added between the
Bcll and Sstl sites using a synthetic oligonucleotide cassette of the form:

5' GATCAAAACATAAAAAACCGGCCTTGGCCCCGGCTTTTTTATTATTTTTGAGCT	3′
3' TTTTGTATTTTTTGGCCGGAACCGGGGCGGCCAAAAAATAATAAAAAC	5′

Sea ID No 1

SstI

designed to contain the B. amyloliquefaciens subtilisin transcriptional terminator (Wells

et al. (1983) Nucleic Acid Research 11:7911-7925).

Site-directed mutagenesis by oligonucleotides used essentially the protocol of Zoller, M. et al. (1983) Meth. Enzymol. 100:468-500: briefly, 5'-phosphorylated oligonucleotide primers were used to introduce the desired mutations on the M13 single-stranded DNA template using the oligonucleotides listed in Table I to substitute for each of the seven methionines found in *B. licheniformis* alpha-amylase. Each mutagenic oligonucleotide also introduced a restriction endonuclease site to use as a screen for the linked mutation.

## TABLE I

# Mutagenic Oligonucleotides for the Substitution of the Methionine Residues in B. licheniformis Alpha-Amylase

5'-T GGG ACG CTG GCG CAG TAC TTT GAA TGG T-3' Scal+	Seq	ID	No	2
5'-TG ATG CAG TAC TTT GAA TGG TAC CTG CCC AAT GA-3' Scal+ Kpnl+	Seq	ID	No	3
5'-GAT TAT TTG TTG TAT GCC GAT ATC GAC TAT GAC CAT-3' $\frac{\text{GAT ATC}}{\text{EcoRV+}}$	Seq	ID	No	4
M256A 5'-CG GGG AAG GAG GCC TTT ACG GTA GCT-3' STUI+	Seq	ID	No	5
M304L 5'-GC GGC TAT GAC TTA AGG AAA TTG C-3'	Seq	ID	No	6
5'-C TAC GGG GAT GCA TAC GGG ACG A-3' NSII+	Seq	ID	No	7
M366Y 5'-C TAC GGG GAT TAC TAC GGG ACC AAG GGA GAC TCC C-3' Styl+	Seq	ID	No	8
M438A 5'-CC GGT GG <u>G</u> GCC AAG CGG GCC TAT GTT GGC CGG CAA A-3' Sfil+	Seç	II	No.	, 9

Bold letter indicate base changes introduced by oligonucleotide.

Codon changes indicated in the form M8A, where methionine (M) at position +8 has been changed to alanine (A).

Underlining indicates restriction endonuclease site introduced by oligonucleotide.

The heteroduplex was used to transfect *E. coli* mutL cells (Kramer et al. (1984) Cell 38:879) and, after plaque-purification, clones were analyzed by restriction analysis of

the RF1's. Positives were confirmed by dideoxy sequencing (Sanger et al. (1977) Proc. Natl. Acad. Sci. U.S.A. 74:5463-5467) and the Pstl-Sstl fragments for each subcloned into an *E. coli* vector, plasmid pA4BL.

# Plasmid pA4BL

Following the methods described in US application 860,468 (Power et al.), which is incorporated herein by reference, a silent PstI site was introduced at codon +1 (the first amino-acid following the signal cleavage site) of the *aprE* gene from pS168-1 (Stahl, M.L. and Ferrari, E. (1984) J. Bacter. 158:411-418). The *aprE* promoter and signal peptide region was then cloned out of a pJH101 plasmid (Ferrari, F.A. et al. (1983) J. Bacter. 154:1513-1515) as a HindIII-PstI fragment and subcloned into the pUC18-derived plasmid JM102 (Ferrari, E. and Hoch, J.A. (1989) Bacillus, ed. C.R. Harwood, Plenum Pub., pp. 57-72). Addition of the PstI-SstI fragment from *B. licheniformis* alpha-amylase gave pA4BL (Fig. 5) having the resulting *aprE* signal peptide-amylase junction as shown in Fig. 6.

## Transformation Into B. subtilis

pA4BL is a plasmid able to replicate in *E. coli* and integrate into the *B. subtilis* chromosome. Plasmids containing different variants were transformed into *B. subtilis* (Anagnostopoulos, C. and Spizizen, J. (1961) J. Bacter. 81:741-746) and integrated into the chromosome at the *aprE* locus by a Campbell-type mechanism (Young, M. (1984) J. Gen. Microbiol. 130:1613-1621). The *Bacillus subtilis* strain BG2473 was a derivative of I168 which had been deleted for amylase (Δ*amyE*) and two proteases (Δ*apr*, Δ*npr*) (Stahl, M.L. and Ferrari, E., J. Bacter. 158:411-418 and US Patent 5,264,366, incorporated herein by reference). After transformation the *sacU*32(Hy) (Henner, D.J. et al. (1988) J. Bacter. 170:296-300) mutation was introduced by PBS-1

mediated transduction (Hoch, J.A. (1983) 154:1513-1515).

GCT-3'

N-terminal analysis of the amylase expressed from pA4BL in *B. subtilis* showed it to be processed having four extra alanines at the N-terminus of the secreted amylase protein ("A4 form"). These extra residues had no significant, deleterious effect on the activity or thermal stability of the A4 form and in some applications may enhance performance. In subsequent experiments the correctly processed forms of the *licheniformis* amylase and the variant M197T were made from a very similar construction (see Fig. 6).

Specifically, the 5' end of the A4 construction was subcloned on an EcoRI-SstII fragment, from pA4BL (Fig. 5) into M13BM20 (Boehringer Mannheim) in order to obtain a coding-strand template for the mutagenic oligonucleotide below:

5'-CAT CAG CGT CCC ATT AAG ATT TGC AGC CTG CGC AGA CAT GTT

Seq ID No 10

This primer eliminated the codons for the extra four N-terminal alanines, correct forms being screened for by the absence of the Pstl site. Subcloning the EcoRl-Sstll fragment back into the pA4BL vector (Fig. 5) gave plasmid pBLapr. The M197T substitution could then be moved, on a Sstll-Sstl fragment, out of pA4BL (M197T) into the complementary pBLapr vector to give plasmid pBLapr (M197T). N-terminal analysis of the amylase expressed from pBLapr in *B. subtilis* showed it to be processed with the same N-terminus found in *B. licheniformis* alpha-amylase.

## Example 2

## Oxidative Sensitivity of Methionine Variants

B. licheniformis alpha-amylase, such as Spezyme® AA20 (commercially available from Genencor International, Inc.), is inactivated rapidly in the presence of hydrogen peroxide (Fig. 7). Various methionine variants were expressed in shake-flask cultures of B. subtilis and the crude supernatants purified by ammonium sulphate cuts. The amylase was precipitated from a 20% saturated ammonium sulphate supernatant by raising the ammonium sulphate to 70% saturated, and then resuspended. The variants were then exposed to 0.88M hydrogen peroxide at pH 5.0, at 25°C. Variants at six of the methionine positions in B. licheniformis alpha-amylase were still subject to oxidation by peroxide while the substitution at position +197 (M197L) showed resistance to peroxide oxidation. (See Fig. 7.) However, subsequent analysis described in further detail below showed that while a variant may be susceptible to oxidation at pH 5.0, 25°C, it may exhibit altered/enhanced properties under different conditions (i.e., liquefaction).

# Example 3

## Construction of All Possible Variants at Position 197

All of the M197 variants (M197X) were produced in the A4 form by cassette mutagenesis, as outlined in Fig. 8:

 Site directed mutagenesis (via primer extension in M13) was used to make M197A using the mutagenic oligonucleotide below:

5'-GAT TAT TTG GCG TAT GCC GAT ATC GAC TAT GAC CAT-3'

ECORV+

ClaiSeq ID No 11

which also inserted an EcoRV site (codons 200-201) to replace the Clal site (codons 201-202).

(codons 201-202).

 Then primer LAAM12 (Table II) was used to introduce another silent restriction site (BstBI) over codons 186-188.

- 3) The resultant M197A (BstBl+, EcoRV+) variant was then subcloned (Pstl-Sstl fragment) into plasmid pA4BL and the resultant plasmid digested with BstBl and EcoRV and the large vector-containing fragment isolated by electroelution from agarose ge!.
- 4) Synthetic primers LAAM14-30 (Table II) were each annealed with the largely complementary common primer LAAM13 (Table II). The resulting cassettes encoded for all the remaining naturally occurring amino acids at position +197 and were ligated, individually, into the vector fragment prepared above.

# TABLE II

# Synthetic Oligonucleotides Used for Cassette Mutagenesis to Produce M197X Variants

LAAM12	GG GAA GT <u>T TCG AA</u> T GAA AAC G	Seq ID No 12
LAAM13	X197bs (Ecorv) $\underline{\text{GTC}}$ GGC at $\underline{\text{A}}$ TG $\underline{\text{CAT}}$ at a atc ata GTT GCC GTT TTC ATT	Seq ID No 13 (BstBI)
LAAM14	1197 (BStBI) CG AAT GAA AAC GGC AAC TAT GAT TAT TTG $\underline{\text{ATC}}$ TAT GCC (	Seq ID No 14 GA <u>C</u> (EcoRV-)
LAAM15	F197 (BSIBI) CG AAT GAA AAC GGC AAC TAT GAT TAT TTG $\overline{\text{TTC}}$ TAT GCC (	Seq ID No 15 SAC (EcoRV-)
LAAM16	V197 (BstBI) CG AAT GAA AAC GGC AAC TAT GAT TAT TTG $\underline{GTT}$ TAT GCC (	Seq ID No 16 GAC (EcoRV-)
LAAM17	S197 (BstBI) CG AAT GAA AAC GGC AAC TAT GAT TAT TTG $\underline{\mathrm{AGC}}$ TAT GCC	Seq ID No 17 GA <u>C</u> (EcoRV-)
LAAM18	P197 (BstBI) CG AAT GAA AAC GGC AAC TAT GAT TAT TTG $\underline{\text{CCT}}$ TAT GCC	Seq ID No 18 GAC (EcoRV-)
LAAM19	T197 (BStBI) CG AAT GAA AAC GGC AAC TAT GAT TAT TTG $\underline{ACA}$ TAT GCC	Seq ID No 19 GAC (EcoRV-)
LAAM20	Y197 (BstBI) CG AAT GAA AAC GGC AAC TAT GAT TAT TTG <u>TAC</u> TAT GCC	Seq ID No 20 GAC (EcoRV-)

WO 94/18314		PCT/US94/01553
LAAM21	H197 (BStBI) CG AAT GAA AAC GGC AAC TAT GAT TAT TTG <u>CAC</u> TAT	Seq ID No 21 FGCC GAC (EcoRV-)
LAAM22	G197 (BstBI) CG AAT GAA AAC GGC AAC TAT GAT TAT TTG $\underline{GGC}$ TAT	Seq ID No 22 F GCC GAC (EcoRV-)
LAAM23	0197 (BstBI) CG AAT GAA AAC GGC AAC TAT GAT TAT TTG <u>CAA</u> TAT	Seq ID No 23 F GCC GAC (EcoRV-)
LAAM24	N197 (BSIBI) CG AAT GAA AAC GGC AAC TAT GAT TAT TTG <u>AAC</u> TAT	Seq ID No 24 F GCC GAC (EcoRV-)
LAAM25	K197 (BstBI) CG AAT GAA AAC GGC AAC TAT GAT TAT TTG <u>AAA</u> TA	Seq ID No 25 F GCC GAC (EcoRV-)
LAAM26	D197 (BstBI) CG AAT GAA AAC GGC AAC TAT GAT TAT TTG <u>GAT</u> TAT	Seq ID No 26 GCC GAC (EcoRV-)
LAAM27	E197 (BstBI) CG AAT GAA AAC GGC AAC TAT GAT TAT TTG <u>GAA</u> TA <sup>*</sup>	Seq ID No 27 F GCC GAC (EcoRV-)
LAAM28	C197 (BstBI) CG AAT GAA AAC GGC AAC TAT GAT TAT TTG <u>TGT</u> TAT	Seq ID No 28 GCC GAC (EcoRV-)
LAAM29	W197 (BstBI) CG AAT GAA AAC GGC AAC TAT GAT TAT TTG <u>TGG</u> TAT	Seq ID No 29 FGCC GAC (EcoRV-)

The cassettes were designed to destroy the EcoRV site upon ligation, thus plasmids from *E. coli* transformants were screened for loss of this unique site. In addition, the common bottom strand of the cassette contained a frame-shift and encoded a Nsil site, thus transformants derived from this strand could be eliminated by screening for the presence of the unique Nsil site and would not be expected, in any case, to lead to expression of active amylase.

(Bs(B)) CG AAT GAA AAC GGC AAC TAT GAT TAT TTG AGA TAT GCC GAC (EcoRV-)

Seq ID No 30

LAAM30

R197

Positives by restriction analysis were confirmed by sequencing and transformed in *B. subtilis* for expression in shake-flask cultures (Fig. 9). The specific activity of certain of the M197X mutants was then determined using a soluble substrate assay. The data generated using the following assay methods are presented below in Table III.

Soluble Substrate Assay: A rate assay was developed based on an end-point assay kit

supplied by Megazyme (Aust.) Pty. Ltd.: Each vial of substrate (p-nitrophenyl maltoheptaoside, BPNPG7) was dissolved in 10ml of sterile water, followed by a 1 to 4 dilution in assay buffer (50mM maleate buffer, pH 6.7, 5mM calcium chloride, 0.002% Tween20). Assays were performed by adding 10µt of amylase to 790µt of the substrate in a cuvette at 25°C. Rates of hydrolysis were measured as the rate of change of absorbance at 410nm, after a delay of 75 seconds. The assay was linear up to rates of 0.4 absorption units/min.

The amylase protein concentration was measured using the standard Bio-Rad assay (Bio-Rad Laboratories) based on the method of Bradford, M. (1976) Anal. Biochem. 72:248) using bovine serum albumin standards.

Starch Hydrolysis Assay: The standard method for assaying the alpha-amylase activity of Spezyme® AA20 was used. This method is described in detail in Example 1 of USSN 07/785,624, incorporated herein by reference. Native starch forms a blue color with iodine but fails to do so when it is hydrolyzed into shorter dextrin molecules. The substrate is soluble Lintner starch 5gm/liter in phosphate buffer, pH 6.2 (42.5gm/liter potassium dihydrogen phosphate, 3.16gm/liter sodium hydroxide). The sample is added in 25mM calcium chloride and activity is measured as the time taken to give a negative iodine test upon incubation at 30°C. Activity is recorded in liquefons per gram or ml (LU) calculated according to the formula:

$$LU/mI \text{ or } LU/g = \underbrace{\phantom{0}570}_{\phantom{0}} \times D$$

Where LU = liquefon unit

V = volume of sample (5ml)

t = dextrinization time (minutes)

D = dilution factor = dilution volume/ml or g of added enzyme.

TABLE III

	SPECIFIC ACTIVITY (as %	of AA20 value) on:
ALPHA-AMYLASE	Soluble Substrate	Starch
3	100	100
Spezyme® AA20		
A4 form	105	115
M15L (A4 form)	93	94
M15L	85	103
M197T (A4 form)	75	83
M197T	62	81
M197A (A4 form)	88	89
M197C	85	85
M197L (A4 form)	51	17

Example 4

## Characterization of Variant M15L

Variant M15L made as per the prior examples did not show increased amylase activity (Table III) and was still inactivated by hydrogen peroxide (Fig. 7). It did, however, show significantly increased performance in jet-liquefaction of starch, especially at low pH as shown in Table IV below.

Starch liquefaction was typically performed using a Hydroheater M 103-M steam jet equipped with a 2.5 liter delay coil behind the mixing chamber and a terminal back pressure valve. Starch was fed to the jet by a Moyno pump and steam was supplied by a 150 psi steam line, reduced to 90-100 psi. Temperature probes were installed just after the Hydroheater jet and just before the back pressure valve.

Starch slurry was obtained from a corn wet miller and used within two days. The starch was diluted to the desired solids level with deionized water and the pH of the starch was adjusted with 2% NaOH or saturated Na<sub>2</sub>CO<sub>3</sub>. Typical liquefaction conditions were:

> Starch 32%-35% solids Calcium

40-50 ppm (30 ppm added) рΗ 5.0-6.0 Alpha-amylase 12-14 LU/g starch dry basis

Starch was introduced into the jet at about 350 ml/min. The jet temperature was held at 105°-107°C. Samples of starch were transferred from the jet cooker to a 95°C second stage liquefaction and held for 90 minutes.

The degree of starch liquefaction was measured immediately after the second stage liquefaction by determining the dextrose equivalence (DE) of the sample and by testing for the presence of raw starch, both according to the methods described in the Standard Analytical Methods of the Member Companies of the Corn Refiners Association, Inc., sixth edition. Starch, when treated generally under the conditions given above and at pH 6.0, will yield a liquefied starch with a DE of about 10 and with no raw starch. Results of starch liquefaction tests using mutants of the present invention are provided in Table IV.

TABLE IV

Performance of

Variants M15L (A4 form) and M15L in Starch Liquefaction

		<u>на</u>	DE after 90 Mins.
Spezyme® M15L (A4 Spezyme® M15L (A4	form) AA20	5.9 5.9 5.2 5.2	9.9 10.4 1.2 2.2
Spezyme®	AA20	5.9	9.3*
M15L		5.9	11.3*
Spezyme®	AA20	5.5	3.25**
M15L		5.5	6.7**
Spezyme®	AA20	5.2	0.7**
M15L		5.2	3.65**

# Example 5

# Construction of M15X Variants

Following generally the processes described in Example 3 above, all variants at M15 (M15X) were produced in native *B. licheniformis* by cassette mutagenesis, as outlined in Fig. 12:

 Site directed mutagenesis (via primer extension in M13) was used to introduce unique restriction sites flanking the M15 codon to facilitate insertion of a mutagenesis cassette. Specifically, a BstB1 site at codons 11-13 and a Msc1 site at codons 18-20 were introduced using the two oligonucleotides shown below.

M15XBstB1 5'-G ATG CAG TAT TTC GAA CTGG TAT A-3' BstB1 Seq ID No 48

M15XMsc1 5'-TG CCC AAT GAT GGC CAA CAT TGG AAG-3' Msc1 Seq ID No 49

<sup>\*</sup> average of three experiments

<sup>\*\*</sup> average of two experiments

2) The vector for M15X cassette mutagenesis was then constructed by subcloning the Sfi1-Sstll fragment from the mutagenized amylase (BstB1+, Msc1+) into plasmid pBLapr. The resulting plasmid was then digested with BstB1 and Msc1 and the large vector fragment isolated by electroelution from a polyacrylamide gel.

3) Mutagenesis cassettes were created as with the M197X variants. Synthetic oligomers, each encoding a substitution at codon 15, were annealed to a common bottom primer. Upon proper ligation of the cassette to the vector, the Msc1 is destroyed allowing for screening of positive transformants by loss of this site. The bottom primer contains an unique SnaB1 site allowing for the transformants derived from the bottom strand to be eliminated by screening for the SnaB1 site. This primer also contains a frameshift which would also eliminate amylase expression for the mutants derived from the common bottom strand.

The synthetic cassettes are listed in Table V and the general cassette mutagenesis strategy is illustrated in Figure 12.

TABLE V

Synthetic Oligonucleotides Used for Cassette Mutagenesis
to Produce M15X Variants

M15A	(BstB1) C GAA	TGG TAT GCT	CCC AAT GAC	GG (Mscl)	seq ID No 50
M15R	(BstB1) C GAA	TGG TAT CGC	CCC AAT GAC	GG (Msc1)	Seq ID No 51
M15N	(BstB1) C GAA	TGG TAT AAT	CCC AAT GAC	GG (Mscl)	Seq ID No 52
M15D	(BstB1) C GAA	TGG TAT GAT	CCC AAT GAC	GG (Msc1)	Seq ID No 53
м15н	(BstB1) C GAA	TGG TAT CAC	CCC AAT GAC	GG (Mscl)	Seq ID No 54
M15K	(BstB1) C GAA	TGG TAT AAI	CCC AAT GAC	GG (Mscl)	Seq ID No 55
M15P	(BstB1) C GAA	TGG TAT CCC	CCC AAT GAC	GG (Msc1)	Seq ID No 56
M15S	(BstB1) C GAA	TGG TAT TC	CCC AAT GAC	GG (Msc1)	Seg ID No 57
M15T	(BstBl) C GAA	TGG TAC AC	CCC AAT GAC	GG (Msc1)	Seq ID No 58
M15V	(BstB1) C GAA	TGG TAT GT	CCC AAT GAC	GG (Msc1)	Seq ID No 59
M15C	(BstB1) C GAA	TGG TAT TG	CCC AAT GAC	GG (Msc1)	Seq ID No 60
M15Q	(BstB1) C GAA	TGG TAT CAL	CCC AAT GAC	GG (Msc1)	Seq ID No 61
M15E	(BstB1) C GAA	TGG TAT GAZ	CCC AAT GAC	GG (Msc1)	Seq ID No 62
M15G	(BstB1) C GAA	TGG TAT GG	CCC AAT GAC	GG (Msc1)	Seq ID No 63
M15I	(BstB1) C GAA	TGG TAT AT	CCC AAT GAC	GG (Msc1)	Seq ID No 64
M15F	(BstB1) C GAA	TGG TAT TT	CCC AAT GAC	GG (Mscl)	Seq ID No 65
M15W	(BstB1) C GAA	TGG TAC TGG	CCC AAT GAC	GG (Msc1)	Seq ID No 66
M15Y	(BstB1) C GAA	TGG TAT TAT	CCC AAT GAC	GG (Msc1)	Seq ID No 67
M15X (bott	(Msc1) CC GTC .om strand)	ATT GGG ACT	r ACG TAC CAT	T (BstB1)	Seq ID No 68

Underline indicates codon changes at amino acid position 15.

Conservative substitutions were made in some cases to prevent introduction of new restriction sites.

## Example 6

## Bench Liquefaction with M15X Variants

Eleven alpha-amylase variants with substitutions for M15 made as per Example 5 were assayed for activity, as compared to Spezyme® AA20 (commercially available from Genencor International, Inc.) in liquefaction at pH 5.5 using a bench liquefaction system. The bench scale liquefaction system consisted of a stainless steel coil (0.25 inch diameter, approximately 350 ml volume) equipped with a 7 inch long static mixing element approximately 12 inches from the anterior end and a 30 psi back pressure valve at the posterior end. The coil, except for each end, was immersed in a glycerol-water bath equipped with thermostatically controlled heating elements that maintained the bath at 105-106°C.

Starch slurry containing enzyme, maintained in suspension by stirring, was introduced into the reaction coil by a piston driven metering pump at about 70 ml/min. The starch was recovered from the end of the coil and was transferred to the secondary hold (95°C for 90 minutes). Immediately after the secondary hold, the DE of the liquefied starch was determined, as described in Example 4. The results are shown in Fig. 16.

## Example 7

# Characterization of M197X Variants

As can be seen in Fig. 9, there was a wide range of amylase activity (measured in the soluble substrate assay) expressed by the M197X (A4 form) variants. The amylases were partially purified from the supernatants by precipitation with two volumes of ethanol and resuspension. They were then screened for thermal stability (Fig. 10) by heating at 95°C for 5 minutes in 10mM acetate buffer pH 5.0, in the presence of 5mM calcium chloride: the A4 wild-type retained 28% of its activity after incubation. For

M197W and M197P we were unable to recover active protein from the supernatants. Upon sequencing, the M197H variant was found to contain a second mutation, N190K. M197L was examined in a separate experiment and was one of the lowest thermally stable variants. There appears to be a broad correlation between expression of amylase activity and thermal stability. The licheniformis amylase is restricted in what residues it can accommodate at position 197 in terms of retaining or enhancing thermal stability: cysteine and threonine are preferred for maximal thermal stability under these conditions whereas alanine and isoleucine are of intermediate stability. However, other substitutions at position +197 result in lowered thermal stability which may be useful for other applications. Additionally, different substitutions at +197 may have other beneficial properties, such as altered pH performance profile or altered oxidative stability. For example, the M197C variant was found to inactivate readily by air oxidation but had enhanced thermal stability. Conversely, compared to the M197L variant, both M197T and M197A retained not only high thermal stability (Fig. 10), but also high activity (Table III), while maintaining resistance to inactivation by peroxide at pH 5 to pH 10 (Fig. 7).

## Example 8

# Stability and Performance in Detergent Formulation

The stability of the M197T (A4 form), M197T and M197A (A4 form) was measured in automatic dish care detergent (ADD) matrices. 2ppm Savinase™ (a protease, commercially available from Novo Industries, of the type commonly used in ADD) were added to two commercially available bleach-containing ADD's: Cascade™ (Procter and Gamble, Ltd.) and Sunlight™ (Unilever) and the time course of inactivation of the amylase variants and Termamyl™ (a thermally stable alpha-amylase available from Novo Nordisk, A/S) followed at 65°C. The concentration of ADD product used in both cases

was equivalent to 'pre-soak' conditions: 14gm product per liter of water (7 grams per gallon hardness). As can be seen (Figs. 11a and 11b), both forms of the M197T variant were much more stable than Termamyl\* and M197A (A4 form), which were inactivated before the first assay could be performed. This stability benefit was seen in the presence or absence of starch as determined by the following protocol. Amylases were added to 5ml of ADD and Savinase\*, prewarmed in a test tube and, after vortexing, activities were assayed as a function of time, using the soluble substrate assay. The "+ starch" tube had spaghetti starch baked onto the sides (140°C, 60 mins.). The results are shown in Figs. 11a and 11b.

# Example 9

## Characterization of M15X Variants

All M15X variants were propagated in *Bacillus subtilis* and the expression level monitored as shown in Fig. 13. The amylase was isolated and partially purified by a 20-70% ammonium sulfate cut. The specific activity of these variants on the soluble substrate was determined as per Example 3 (Fig. 14). Many of the M15X amylases have specific activities greater than that of Spezyme® AA20. A benchtop heat stability assay was performed on the variants by heating the amylase at 90°C for 5 min. in 50 mM acetate buffer pH 5 in the presence of 5 mM CaCl<sub>2</sub> (Fig. 15). Most of the variants performed as well as Spezyme® AA20 in this assay. Those variants that exhibited reasonable stability in this assay (reasonable stability defined as those that retained at least about 60% of Spezyme® AA20's heat stability) were tested for specific activity on starch and for liquefaction performance at pH 5.5. The most interesting of those mutants are shown in Fig. 16. M15D, N and T, along with L, outperformed Spezyme® AA20 in liquefaction at pH 5.5 and have increased specific activities in both the soluble substrate and starch hydrolysis assays.

Generally, we have found that by substituting for the methionine at position 15, we can provide variants with increased low pH-liquefaction performance and/or increased specific activity.

## Example 10

## Tryptophan Sensitivity to Oxidation

Chloramine-T (sodium N-chloro-*p*-toluenesulfonimide) is a selective oxidant, which oxidizes methionine to methionine sulfoxide at neutral or alkaline pH. At acidic pH, chloramine-T will modify both methionine and tryptophan (Schechter, Y., Burstein, Y. and Patchornik, A. (1975) Biochemistry 14 (20) 4497-4503). Fig. 17 shows the inactivation of *B. licheniformis* alpha-amylase with chloramine-T at pH 8.0 (AA20 = 0.65 mg/ml, M197A = 1.7 mg/ml, M197L = 1.7 mg/ml). The data shows that by changing the methionine at position 197 to leucine or alanine, the inactivation of alpha-amylase can be prevented. Conversely, as shown in Fig. 18, at pH 4.0 inactivation of the M197A and M197L proceeds, but require more equivalents of chloramine-T (Fig. 18; AA20 = 0.22 mg/ml, M197A = 4.3 mg/ml, M197L = 0.53 mg/ml; 200 mM NaAcetate at pH 4.0). This suggests that a tryptophan residue is also implicated in the chloramine-T mediated inactivation event. Furthermore, tryptic mapping and subsequent amino acid sequencing indicated that the tryptophan at position 138 was oxidized by chloramine-T (data not shown). To prove this, site-directed mutants were made at tryptophan 138 as provided below:

## Preparation of Alpha-Amylase Double Mutants W138 and M197

Certain variants of W138 (F, Y and A) were made as double mutants, with M197T (made as per the disclosure of Example 3). The double mutants were made following the methods described in Examples 1 and 3. Generally, single negative strands of DNA

were prepared from an M13MP18 clone of the 1.72kb coding sequence (Pst I-Sst I) of the *B. licheniformis* alpha-amylase M197T mutant. Site-directed mutagenesis was done using the primers listed below, essentially by the method of Zoller, M. et al. (1983) except T4 gene 32 protein and T4 polymerase were substituted for klenow. The primers all contained unique sites, as well as the desired mutation, in order to identify those clones with the appropriate mutation.

#### Tryptophan 138 to Phenylalanine

133 134 135 136 137 138 139 140 141 142 143
CAC CTA ATT AAA GCT TTC ACA CAT TTT CAT TTT Seq ID No 42
Hind III

#### Tryptophan 138 to Tyrosine

133 134 135 136 137 138 139 140 141 142 143 CAC CTA ATT AAA GCT TAC ACA CAT TTT CAT TTT Hind III

Seq ID No 43

Tryptophan 138 to Alanine - This primer also engineers unique sites upstream and downstream of the 138 position.

127 128 129 130 131 132 133 134 135 136 137 138 139 140 141 142 C CGC GTA ATT  $\frac{TCC}{BS}$  GAA CAC CTA ATT AAA GCC GCA ACA CAT TTT CAT BSpE I

143 144 145 146 147 TTT <u>CCC GGG</u> CGC GGC AG Xma I

Sea ID No 44

Mutants were identified by restriction analysis and W138F and W138Y confirmed by DNA sequencing. The W138A sequence revealed a nucleotide deletion between the unique BspE I and Xma I sites, however, the rest of the gene sequenced correctly. The 1.37kb SstII/SstI fragment containing both W138X and M197T mutations was moved from M13MP18 into the expression vector pBLapr resulting in pBLapr (W138F, M197T) and pBLapr (W138Y, M197T). The fragment containing unique BspE I and Xma I sites was cloned into pBLapr (BspE I, Xma I, M197T) since it is useful for cloning cassettes containing other amino acid substitutions at position 138.

#### Single Mutations at Amino Acid Position138

Following the general methods described in the prior examples, certain single variants of W138 (F. Y. L. H and C) were made.

The 1.24kb Asp718-Sstl fragment containing the M197T mutation in plasmid pBLapr (W138X, M197T) of Example 7 was replaced by the wild-type fragment with methionine at 197, resulting in pBLapr (W138F), pBLapr (W138Y) and pBLapr (BspE I, Xma I).

The mutants W138L, W138H and W138C were made by ligating synthetic cassettes into the pBLapr (BspE I, Xma I) vector using the following primers:

### Tryptophan 138 to Leucine

CC GGA GAA CAC CTA ATT AAA GCC CTA ACA CAT TTT CAT TTT C
Seq ID No 45

#### Tryptophan 138 to Histidine

CC GGA GAA CAC CTA ATT AAA GCC CAC ACA CAT TTT CAT TTT C
Seq ID No 46

# Tryptophan 138 to Cysteine

CC GGA GAA CAC CTA ATT AAA GCC TGC ACA CAT TTT CAT TTT C

Seq ID No 47

Reaction of the double mutants M197T/W138F and M197T/W138Y with chloramine-T was compared with wild-type (AA20 = 0.75 mg/ml, M197T/W138F = 0.64 mg/ml, M197T/W138Y = 0.60 mg/ml; 50 mM NaAcetate at pH 5.0). The results shown in Fig. 19 show that mutagenesis of tryptophan 138 has caused the variant to be more resistant to chloramine-T.

#### SEQUENCE LISTING

- (1) GENERAL INFORMATION:
  - (i) APPLICANT: GENENCOR INTERNTIONAL, INC.
  - (ii) TITLE OF INVENTION: Oxidatively Stable Alpha-Amylase
  - (iii) NUMBER OF SEQUENCES: 68
  - (iv) CORRESPONDENCE ADDRESS:
    - (A) ADDRESSEE: Genencor International, Inc.
    - (B) STREET: 180 Kimball Way
    - (C) CITY: South San Francisco
    - (D) STATE: CA (E) COUNTRY: USA
    - (F) ZIP: 94080
    - (v) COMPUTER READABLE FORM:
      - (A) MEDIUM TYPE: Floppy disk
        - (B) COMPUTER: IBM PC compatible (C) OPERATING SYSTEM: PC-DOS/MS-DOS
        - (D) SOFTWARE: PatentIn Release #1.0, Version #1.25
  - (vi) CURRENT APPLICATION DATA:
    - (A) APPLICATION NUMBER:
    - (B) FILING DATE: (C) CLASSIFICATION:
  - (viii) ATTORNEY/AGENT INFORMATION: (A) NAME: Horn, Margaret A.
    - (B) REGISTRATION NUMBER: 33,401
    - (C) REFERENCE/DOCKET NUMBER: GC220-2
    - (ix) TELECOMMUNICATION INFORMATION:
      - (A) TELEPHONE: (415) 742-7536 (B) TELEFAX: (415) 742-7217
- (2) INFORMATION FOR SEQ ID NO:1:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 56 base pairs
    - (B) TYPE: nucleic acid
    - (C) STRANDEDNESS: single (D) TOPOLOGY: linear
  - (ii) MOLECULE TYPE: DNA (genomic)
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:1:
- GATCAAAACA TAAAAAACCG GCCTTGGCCC CGCCGGTTTT TTATTATTTT TGAGCT

- (2) INFORMATION FOR SEQ ID NO:2:
  - (i) SEQUENCE CHARACTERISTICS:

		(A) LENGTH: 29 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
	(ii)	MOLECULE TYPE: DNA (genomic)	
	(xi)	SEQUENCE DESCRIPTION: SEQ ID NO:2:	
TGG	ACGC	TG GCGCAGTACT TTGAATGGT	29
(2)	INFO	RMATION FOR SEQ ID NO:3:	
	(i)	SEQUENCE CHARACTERISTICS: (A) LENGTH: 34 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOCY: linear	
	(ii)	MOLECULE TYPE: DNA (genomic)	
	(xi)	SEQUENCE DESCRIPTION: SEQ ID NO:3:	
TGA:	rgcag	TA CTTTGAATGG TACCTGCCCA ATGA	34
(2)	INFO	RMATION FOR SEQ ID NO:4:	
	(i)	SEQUENCE CHARACTERISTICS: (A) LENGTH: 36 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOFOLOGY: linear	
	(ii)	MOLECULE TYPE: DNA (genomic)	
	(xi)	SEQUENCE DESCRIPTION: SEQ ID NO:4:	
GAT'	PATTT	GT TGTATGCCGA TATCGACTAT GACCAT	36
(2)	INFO	RMATION FOR SEQ ID NO:5:	
	(i)	SEQUENCE CHARACTERISTICS: (A) LENGTH: 26 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
	(ii)	MOLECULE TYPE: DNA (genomic)	
	(xi)	SEQUENCE DESCRIPTION: SEQ ID NO:5:	
CGG	GGAAG	GA GGCCTTTACG GTAGCT	26
(2)	INFO	RMATION FOR SEQ ID NO:6:	
	(i)	SEQUENCE CHARACTERISTICS: (A) LENGTH: 24 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	

(ii) MOLECULE TYPE: DNA (genomic)	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:6:	
GCGGCTATGA CTTAAGGAAA TTGC	24
(2) INFORMATION FOR SEQ ID NO:7:	
(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 23 hase pairs (B) TYPE: nucleic acid (C) STRANDEDMESS: single (D) TOFOLOGY: linear	
(ii) MOLECULE TYPE: DNA (genomic)	
(xi) SEQUENCE DESCRIPTION: SEQ ID No:7:	
CTACGGGGAT GCATACGGGA CGA	23
(2) INFORMATION FOR SEQ ID NO:8:	
(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 35 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
(ii) MOLECULE TYPE: DNA (genomic)	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:8:	
CTACGGGGAT TACTACGGGA CCAAGGGAGA CTCCC	35
(2) INFORMATION FOR SEQ ID NO:9:	
(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 36 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
(ii) MOLECULE TYPE: DNA (genomic)	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:9:	
CCGGTGGGGC CAAGCGGGCC TATGTTGGCC GGCAAA	36
(2) INFORMATION FOR SEQ ID NO:10:	
(i) SEQUENCE CHARACTERISTICS: (A) LENGTH 45 base pairs (B) TYPE: nucleic acid (C) STRANEDNESS: single (D) TOPOLOGY: linear	
(ii) MOLECULE TYPE: DNA (genomic)	

(XI) SEQUENCE DESCRIPTION. SEQ IS NO. 10.	
CATCAGCGTC CCATTAAGAT TTGCAGCCTG CGCAGACATG TTGCT	45
(2) INFORMATION FOR SEQ ID NO:11:	
(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 36 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
(ii) MOLECULE TYPE: DNA (genomic)	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:11:	
GATTATTTGG CGTATGCCGA TATCGACTAT GACCAT	36
(2) INFORMATION FOR SEQ ID NO:12:	
(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 21 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
(ii) MOLECULE TYPE: DNA (genomic)	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:12:	
GGGAAGTTTC GAATGAAAAC G	21
(2) INFORMATION FOR SEQ ID NO:13:	
(i) SEQUENCE CHARACTERISTICS:  (A) LENGTH: 36 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
(ii) MOLECULE TYPE: DNA (genomic)	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:13:	
GTCGGCATAT GCATATAATC ATAGTTGCCG TTTTCATT	38
(2) INFORMATION FOR SEQ ID NO:14:	
(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 41 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
(ii) MOLECULE TYPE: DNA (genomic)	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:14:	
CGAATGAAAA CGGCAACTAT GATTATTTGA TCTATGCCGA C	41

(2) INFORMATION FOR SEQ ID NO:15:	
(i) SEQUENCE CHARACTERISTICS:  (A) LENCTH: 41 base pairs  (B) TYPE: nucleic acid  (C) STRANDEDMESS: single  (D) TOPOLOGY: linear	
(ii) MOLECULE TYPE: DNA (genomic)	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:15:	
CGAATGAAAA CGGCAACTAT GATTATTTGT TCTATGCCGA C	41
(2) INFORMATION FOR SEQ ID NO:16:	
(i) SEQUENCE CHARACTERISTICS:  (A) LENGTH: 41 base pairs  (B) TYPE: nucleic acid  (C) STRANDEDNESS: single  (D) TOPOLOGY: linear	
(ii) MOLECULE TYPE: DNA (genomic)	
•	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:16:	
CGAATGAAAA CGGCAACTAT GATTATTTGG TTTATGCCGA C	41
(2) INFORMATION FOR SEQ ID NO:17:	
(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 41 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPGLOOF: linear	
(ii) MOLECULE TYPE: DNA (genomic)	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:17:	
CGAATGAAAA CGGCAACTAT GATTATTTGA GCTATGCCGA C	41
(2) INFORMATION FOR SEQ ID NO:18:	
(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 41 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
(ii) MOLECULE TYPE: DNA (genomic)	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:18:	
CGAATGAAAA CGGCAACTAT GATTATTTGC CTTATGCCGA C	41
	41
(2) INFORMATION FOR SEQ ID NO:19:	
(i) SEQUENCE CHARACTERISTICS:	

(A) LENGTH: 41 base pairs (B) TYPE: nucleic acid (C) STRANDEDWESS: single (D) TOPOLOGY: linear	
(ii) MOLECULE TYPE: DNA (genomic)	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:19:	
CGAATGAAAA CGGCAACTAT GATTATTTGA CATATGCCGA C	41
(2) INFORMATION FOR SEQ ID NO:20:	
(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 41 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
(ii) MOLECULE TYPE: DNA (genomic)	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:20:	
CGAATGAAAA CGGCAACTAT GATTATTTGT ACTATGCCGA C	41
(2) INFORMATION FOR SEQ ID NO:21:	
(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 41 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
(ii) MOLECULE TYPE: DNA (genomic)	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:21:	
CGAATGAAAA CGGCAACTAT GATTATTTGC ACTATGCCGA C	41
(2) INFORMATION FOR SEQ ID NO:22:	
(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 41 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
(ii) MOLECULE TYPE: DNA (genomic)	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:22:	
CGAATGAAAA CGGCAACTAT GATTATTTGG GCTATGCCGA C	41
(2) INFORMATION FOR SEQ ID NO:23:	
(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 41 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	

(ii) MOLECULE TYPE: DNA (genomic)

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:23:	
CGAATGAAAA CGGCAACTAT GATTATTTGC AATATGCCGA C	41
(2) INFORMATION FOR SEQ ID NO:24:	
(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 41 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
(ii) MOLECULE TYPE: DNA (genomic)	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:24:	
CGAATGAAAA CGGCAACTAT GATTATTTGA ACTATGCCGA C	41
(2) INFORMATION FOR SEQ ID NO:25:	
(i) SEQUENCE CHARACTERISTICS: (A) LENOTH: 41 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
(ii) MOLECULE TYPE: DNA (genomic)	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:25:	
GCAATGAAAA CGGCAACTAT GATTATTTGA AATATGCCGA C	41
(2) INFORMATION FOR SEQ ID NO:26:	7.
(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 41 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
(ii) MOLECULE TYPE: DNA (genomic)	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:26:	
CGAATGAAAA CGGCAACTAT GATTATTTGG ATTATGCCGA C	41
(2) INFORMATION FOR SEQ ID NO:27:	
(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 41 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
(ii) MOLECULE TYPE: DNA (genomic)	

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:27: CGAATGAAAA CGGCAACTAT GATTATTTGG AATATGCCGA C 41 (2) INFORMATION FOR SEQ ID NO:28: (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 41 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear (ii) MOLECULE TYPE: DNA (genomic) (xi) SEQUENCE DESCRIPTION: SEQ ID NO:28: CGAATGAAAA CGGCAACTAT GATTATTTGT GTATTGCCGA C 41 (2) INFORMATION FOR SEQ ID NO:29: (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 41 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear (ii) MOLECULE TYPE: DNA (genomic) (xi) SEQUENCE DESCRIPTION: SEQ ID NO:29: CGAATGAAAA CGGCAACTAT GATTATTTGT GGTATGCCGA C 41 (2) INFORMATION FOR SEQ ID NO:30: (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 41 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear (ii) MOLECULE TYPE: DNA (genomic) (xi) SEQUENCE DESCRIPTION: SEQ ID NO:30: CGAATGAAAA CGGCAACTAT GATTATTTGA GATATGCCGA C 41 (2) INFORMATION FOR SEQ ID NO:31: (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 1968 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear (ii) MOLECULE TYPE: DNA (genomic) (xi) SEQUENCE DESCRIPTION: SEQ ID NO:31:

44

AGCTTGAAGA AGTGAAGAAG CAGAGAGGCT ATTGAATAAA TGAGTAGAAA GCGCCATATC 60

GGCGCTTTT	C TTTTGGAAG	A AAATATAGG	AAAATGGTAC	TTGTTAAAA	TTCGGAATAT	120
TTATACAAC	A TCATATGTT	CACATTGAA	A GGGGAGGAGA	ATCATGAAA	AACAAAAACG	180
GCTTTACGC	C CGATTGCTG	CGCTGTTATT	TGCGCTCATC	TTCTTGCTG	CTCATTCTGC	240
AGCAGCGGC	G GCAAATCTT#	ATGGGACGCT	GATGCAGTAT	TTTGAATGGT	ACATGCCCAA	300
TGACGGCCA	A CATTGGAAGO	GTTTGCAAA	CGACTCGGCA	TATTTGGCTG	AACACGGTAT	360
TACTGCCGT	TGGATTCCCC	CGGCATATAA	GGGAACGAGC	CAAGCGGATG	TGGGCTACGG	420
TGCTTACGAC	CTTTATGATT	TAGGGGAGTT	TCATCAAAAA	GGGACGGTTC	GGACAAAGTA	480
CGGCACAAA	GGAGAGCTGC	AATCTGCGAT	CAAAAGTCTT	CATTCCCGCG	ACATTAACGT	540
TTACGGGGAT	GTGGTCATCA	ACCACAAAGG	CGGCGCTGAT	GCGACCGAAG	ATGTAACCGC	600
GGTTGAAGTC	GATCCCGCTG	ACCGCAACCG	CGTAATTTCA	GGAGAACACC	TAATTAAAGC	660
CTGGACACAT	TTTCATTTTC	ceeecece	CAGCACATAC	AGCGATTTTA	AATGGCATTG	720
GTACCATTTT	GACGGAACCG	ATTGGGACGA	GTCCCGAAAG	CTGAACCGCA	TCTATAAGTT	780
TCAAGGAAAG	GCTTGGGATT	GGGAAGTTTC	CAATGAAAAC	GGCAACTATG	ATTATTTGAT	840
GTATGCCGAC	ATCGATTATG	ACCATCCTGA	TGTCGCAGCA	GAAATTAAGA	GATGGGGCAC	900
TTGGTATGCC	AATGAACTGC	AATTGGACGG	TTTCCGTCTT	GATGCTGTCA	AACACATTAA	960
ATTTTCTTTT	TTGCGGGATT	GGGTTAATCA	TGTCAGGGAA	AAAACGGGGA	AGGAAATGTT	1020
TACGGTAGCT	GAATATTGGC	AGAATGACTT	GGGCGCGCTG	GAAAACTATT	TGAACAAAAC	1080
AAATTTTAAT	CATTCAGTGT	TTGACGTGCC	GCTTCATTAT	CAGTTCCATG	CTGCATCGAC	1140
ACAGGGAGGC	GGCTATGATA	TGAGGAAATT	GCTGAACGGT	ACGGTCGTTT	CCAAGCATCC	1200
GTTGAAATCG	GTTACATTTG	TCGATAACCA	TGATACACAG	CCGGGGCAAT	CGCTTGAGTC	1260
GACTGTCCAA	ACATGGTTTA	AGCCGCTTGC	TTACGCTTTT	ATTCTCACAA	GGGAATCTGG	1320
ATACCCTCAG	GTTTTCTACG	GGGATATGTA	CGGGACGAAA	GGAGACTCCC	AGCGCGAAAT	1380
TCCTGCCTTG	AAACACAAAA	TTGAACCGAT	CTTAAAAGCG	AGAAAACAGT	ATGCGTACGG	1440
AGCACAGCAT	GATTATTTCG	ACCACCATGA	CATTGTCGGC	TGGACAAGGG	AAGGCGACAG	1500
CTCGGTTGCA	AATTCAGGTT	TGGCGGCATT	AATAACAGAC	GGACCCGGTG	GGGCAAAGCG	1560
AATGTATGTC	GGCCGGCAAA	ACGCCGGTGA	GACATGGCAT	GACATTACCG	GAAACCGTTC	1620
GGAGCCGGTT	GTCATCAATT	CGGAAGGCTG	GGGAGAGTTT	CACGTAAACG	GCGGGTCGGT	1680
TTCAATTTAT	GTTCAAAGAT	AGAAGAGCAG	AGAGGACGGA	TTTCCTGAAG	GAAATCCGTT	1740
TTTTTATTTT	GCCCGTCTTA	TAAATTTCTT	TGATTACATT	TTATAATTAA	TTTTAACAAA	1800
GTGTCATCAG	CCCTCAGGAA	GGACTTGCTG	ACAGTTTGAA	TCGCATAGGT	AAGGCGGGGA	1860
TGAAATGGCA	ACGTTATCTG	ATGTAGCAAA	GAAAGCAAAT	GTGTCGAAAA	TGACGGTATC	1920
GCGGGTGATC	AATCATCCTG	AGACTGTGAC	GGATGAATTG	AAAAAGCT		1968

<sup>(2)</sup> INFORMATION FOR SEQ ID NO:32:

<sup>(</sup>i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 483 amino acids

- (B) TYPE: amino acid(C) STRANDEDNESS: single
- (C) STRANDEDNESS: sings (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: protein

#### (xi) SEQUENCE DESCRIPTION: SEQ ID NO:32:

Ala Asn Leu Asn Gly Thr Leu Met Gln Tyr Phe Glu Trp Tyr Met Pro Asn Asp Gly Gln His Trp Lys Arg Leu Gln Asn Asp Ser Ala Tyr Leu 20 25 30 Ala Glu His Gly Ile Thr Ala Val Trp Ile Pro Pro Ala Tyr Lys Gly
35 40 45 Thr Ser Gln Ala Asp Val Gly Tyr Gly Ala Tyr Asp Leu Tyr Asp Leu 50 60 Gly Glu Phe His Gln Lys Gly Thr Val Arg Thr Lys Tyr Gly Thr Lys 65 70 75 80 Gly Glu Leu Gln Ser Ala Ile Lys Ser Leu His Ser Arg Asp Ile Asn 85 90 95 Val Tyr Gly Asp Val Val Ile Asn His Lys Gly Gly Ala Asp Ala Thr Glu Asp Val Thr Ala Val Glu Val Asp Pro Ala Asp Arg Asn Arg Val 115 120 125Ile Ser Gly Glu His Leu Ile Lys Ala Trp Thr His Phe His Phe Pro 130 135 140 Gly Arg Gly Ser Thr Tyr Ser Asp Phe Lys Trp His Trp Tyr His Phe 145 150 155 160 Asp Gly Thr Asp Trp Asp Glu Ser Arg Lys Leu Asn Arg Ile Tyr Lys 165 170 175 Phe Gln Gly Lys Ala Trp Asp Trp Glu Val Ser Asn Glu Asn Gly Asn 180 185 190Tyr Asp Tyr Leu Met Tyr Ala Asp Ile Asp Tyr Asp His Pro Asp Val Ala Ala Glu Ile Lys Arg Trp Gly Thr Trp Tyr Ala Asn Glu Leu Gln 210 225 220 Leu Asp Gly Phe Arg Leu Asp Ala Val Lys His Ile Lys Phe Ser Phe 225 230 235 240 Leu Arg Asp Trp Val Asn His Val Arg Glu Lys Thr Gly Lys Glu Met 245 250 255 Phe Thr Val Ala Glu Tyr Trp Gln Asn Asp Leu Gly Ala Leu Glu Asn 260 265 270 Tyr Leu Asn Lys Thr Asn Phe Asn His Ser Val Phe Asp Val Pro Leu 275 280 285 His Tyr Gln Phe His Ala Ala Ser Thr Gln Gly Gly Gly Tyr Asp Met Arg Lys Leu Leu Asn Gly Thr Val Val Ser Lys His Pro Leu Lys Ser

310 315 305 Val Thr Phe Val Asp Asn His Asp Thr Gln Pro Gly Gln Ser Leu Glu 325 330 Ser Thr Val Gln Thr Trp Phe Lys Pro Leu Ala Tyr Ala Phe Ile Leu Thr Arg Glu Ser Gly Tyr Pro Gln Val Phe Tyr Gly Asp Met Tyr Gly 360 Thr Lys Gly Asp Ser Gln Arg Glu Ile Pro Ala Leu Lys His Lys Ile 370 375 380 Glu Pro Ile Leu Lys Ala Arg Lys Gln Tyr Ala Tyr Gly Ala Gln His Asp Tyr Phe Asp His His Asp Ile Val Gly Trp Thr Arg Glu Gly Asp Ser Ser Val Ala Asn Ser Gly Leu Ala Ala Leu Ile Thr Asp Gly Pro Gly Gly Ala Lys Arg Met Tyr Val Gly Arg Gln Asn Ala Gly Glu Thr 435 440 445 Trp His Asp Ile Thr Gly Asn Arg Ser Glu Pro Val Val Ile Asn Ser Glu Gly Trp Gly Glu Phe His Val Asn Gly Gly Ser Val Ser Ile Tyr

(2) INFORMATION FOR SEQ ID NO:33:

Val Gln Arg

- (i) SEQUENCE CHARACTERISTICS:
  - (A) LENGTH: 511 amino acids
  - (B) TYPE: amino acid (C) STRANDEDNESS: single
  - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: protein
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:33:

Met Lys Gln Gln Lys Arg Leu Tyr Ala Arg Leu Leu Thr Leu Leu Phe 1 10 15

Ala Leu Ile Phe Leu Leu Pro His Ser Ala Ala Ala Ala Ala Asn Leu 20 25 30

Asn Gly Thr Leu Met Gln Tyr Phe Glu Trp Tyr Met Pro Asn Asp Gly 35 40 45

His Trp Lys Arg Leu Gln Asn Asp Ser Ala Tyr Leu Ala Glu His Gly

Ile Thr Ala Val Trp Ile Pro Pro Ala Tyr Lys Gly Thr Ser Gln Ala 65 70 75 80

Asp Val Gly Tyr Gly Ala Tyr Asp Leu Tyr Asp Leu Gly Glu Phe His 85 90 95

Gln Lys Gly Thr Val Arg Thr Lys Tyr Gly Thr Lys Gly Glu Leu Gln

			100					105					110		
Ser A		le 15	Lys	Ser	Leu	His	Ser 120	Arg	Asp	Ile	Asn	Val 125	Tyr	Gly	Asp
Val V	al I 30	le	Asn	His	Lys	Gly 135	Gly	Ala	Asp	Ala	Thr 140	Glu	Asp	Val	Thr
Ala V 145	al G	lu	Val	Asp	Pro 150	Ala	Asp	Arg	Asn	Arg 155	Val	Ile	Ser	Gly	Glu 160
His L	eu I	le	Lys	Ala 1 <b>6</b> 5	Trp	Thr	His	Phe	His 170	Phe	Pro	Gly	Arg	Gly 175	Ser
Thr T	yr S	er	Asp 180	Phe	Lys	Trp	His	Trp 185	Tyr	His	Phe	Asp	Gly 190	Thr	Asp
Trp A	sp G	1u 95	Ser	Arg	Lys	Leu	Asn 200	Arg	Ile	Tyr	Lys	Phe 205	Gln	Gly	Lys
Ala T 2	rp A	.sp	Trp	Glu	Val	Ser 215	Asn	Glu	Asn	Gly	Asn 220	Tyr	Asp	Tyr	Leu
Met T 225	yr A	la	Asp	Ile	Asp 230	Tyr	Asp	His	Pro	Asp 235	Val	Ala	Ala	Glu	11e 240
Lys A	rg T	rp	Gly	Thr 245	Trp	Tyr	Ala	Asn	Glu 250	Leu	Gln	Leu	Asp	Gly 255	Phe
Arg L	eu A	sp	Ala 260	Val	Lys	His	Ile	Lys 265	Phe	Ser	Phe	Leu	Arg 270	Asp	Trp
Val A	sn H 2	1 <b>s</b> 75	Val	Arg	Glu	Lys	Thr 280	Gly	Lys	Glu	Met	Phe 285	Thr	Val	Ala
Glu T	yr T 90	rp	Gln	Asn	Asp	Leu 295	Gly	Ala	Leu	Glu	Asn 300	Tyr	Leu	Asn	Lys
Thr A	sn P	he	Asn	His	Ser 310	Val	Phe	Asp	Val	Pro 315	Leu	His	Tyr	Gln	Phe 320
His A	la A	la	Ser	Thr 325	Gln	Gly	Gly	Gly	Tyr 330	Asp	Met	Arg	Lys	Leu 335	Leu
Asn G	ly T		Val 340	Val	Ser	Lys	His	Pro 345	Leu	Lys	Ser	Val	Thr 350	Phe	Val
Asp A	3	55	_				360					365			
	rp P 70	he	Lys	Pro	Leu	Ala 375	Tyr	Ala	Phe	Ile	Leu 380	Thr	Arg	Glu	Ser
Gly T 385	-				390					395					400
Ser G	ln A	rg	Glu	11e 405	Pro	Ala	Leu	Lys	His 410	Lys	Ile	Glu	Pro	11e 415	Leu
Lys A			420					425					430		
His H	4	35					440					445			
Asn S	er G 50	ly:	Leu	Ala	Ala	Leu 455	Ile	Thr	Asp	Gly	Pro 460	Gly	Gly	Ala	Lys

Arg Met Tyr Val Gly Arg Gln Asn Ala Gly Glu Thr Trp His Asp Ile
485

Thr Gly Asn Arg Ser Glu Pro Val Val Ile Asn Ser Glu Gly Trp Gly
495

Glu Phe His Val Asn Gly Gly Ser Val Ser Ile Tyr Val Gln Arg

- (2) INFORMATION FOR SEQ ID NO:34:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 520 amino acids
      - (B) TYPE: amino acid
    - (C) STRANDEDNESS: single
      (D) TOPOLOGY: linear
  - (ii) MOLECULE TYPE: protein

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:34: Met Arg Gly Arg Gly Asn Met Ile Gln Lys Arg Lys Arg Thr Val Ser Phe Arg Leu Val Leu Met Cys Thr Leu Leu Phe Val Ser Leu Pro Ile 20 25 30 Thr Lys Thr Ser Ala Val Asn Gly Thr Leu Met Gln Tyr Phe Glu Trp Tyr Thr Pro Asn Asp Gly Gln His Trp Lys Arg Leu Gln Asn Asp Ala 50 60 Glu His Leu Ser Asp Ile Gly Ile Thr Ala Val Trp Ile Pro Pro Ala 65 70 75 80 Tyr Lys Gly Leu Ser Gln Ser Asp Asn Gly Tyr Gly Pro Tyr Asp Leu 85 90 95 Tyr Asp Leu Gly Glu Phe Gln Gln Lys Gly Thr Val Arg Thr Lys Tyr 100 105 110 Gly Thr Lys Ser Glu Leu Gln Asp Ala Ile Gly Ser Leu His Ser Arg 115 120 125 Asn Val Gln Val Tyr Gly Asp Val Val Leu Asn His Lys Ala Gly Ala Asp Ala Thr Glu Asp Val Thr Ala Val Glu Val Asn Pro Ala Asn Arg 145 150 155 160 Asn Gln Glu Thr Ser Glu Glu Tyr Gln Ile Lys Ala Trp Thr Asp Phe Arg Phe Pro Gly Arg Gly Asn Thr Tyr Ser Asp Phe Lys Trp His Trp 180 185 190 Tyr His Phe Asp Gly Ala Asp Trp Asp Glu Ser Arg Lys Ile Ser Arg Ile Phe Lys Phe Arg Gly Glu Gly Lys Ala Trp Asp Trp Glu Val Ser 210 215 220 Ser Glu Asn Gly Asn Tyr Asp Tyr Leu Met Tyr Ala Asp Val Asp Tyr 225 230 235 240

Asp His Pro Asp Val Val Ala Glu Thr Lys Lys Trp Gly Ile Trp Tyr Ala Asn Glu Leu Ser Leu Asp Gly Phe Arg Ile Asp Ala Ala Lys His Ile Lys Phe Ser Phe Leu Arg Asp Trp Val Gln Ala Val Arg Gln Ala 275 280 285 Thr Gly Lys Glu Met Phe Thr Val Ala Glu Tyr Trp Gln Asn Asn Ala Gly Lys Leu Glu Asn Tyr Leu Asn Lys Thr Ser Phe Asn Gln Ser Val 305 310 . 315 320 Phe Asp Val Pro Leu His Phe Asn Leu Gln Ala Ala Ser Ser Gln Gly 325 330 335 Gly Gly Tyr Asp Met Arg Arg Leu Leu Asp Gly Thr Val Val Ser Arg His Pro Glu Lys Ala Val Thr Phe Val Glu Asn His Asp Thr Gln Pro Gly Gln Ser Leu Glu Ser Thr Val Gln Thr Trp Phe Lys Pro Leu Ala Tyr Ala Phe Ile Leu Thr Arg Glu Ser Gly Tyr Pro Gln Val Phe Tyr 385 390 395 400 Gly Asp Met Tyr Gly Thr Lys Gly Thr Ser Pro Lys Glu Ile Pro Ser 405 410 415 Leu Lys Asp Asn Ile Glu Pro Ile Leu Lys Ala Arg Lys Glu Tyr Ala Tyr Gly Pro Gln His Asp Tyr Ile Asp His Pro Asp Val Ile Gly Trp 435 440 Thr Arg Glu Gly Asp Ser Ser Ala Ala Lys Ser Gly Leu Ala Ala Leu
450 460 Ile Thr Asp Gly Pro Gly Gly Ser Lys Arg Met Tyr Ala Gly Leu Lys 465 470 475 480 Asn Ala Gly Glu Thr Trp Tyr Asp Ile Thr Gly Asn Arg Ser Asp Thr 485 490 495 Val Lys Ile Gly Ser Asp Gly Trp Gly Glu Phe His Val Asn Asp Gly 500 505 510 Ser Val Ser Ile Tyr Val Gln Lys

#### (2) INFORMATION FOR SEQ ID NO:35:

- (i) SEQUENCE CHARACTERISTICS:
  - (A) LENGTH: 548 amino acids
  - (B) TYPE: amino acid
  - (C) STRANDEDNESS: single (D) TOPOLOGY: linear
  - (D) TOPOLOGI. IIMERI
- (ii) MOLECULE TYPE: protein
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:35:

Val Leu Thr Phe His Arg Ile Ile Arg Lys Gly Trp Met Phe Leu Leu Ala Phe Leu Leu Thr Ala Ser Leu Phe Cys Pro Thr Gly Arg His Ala 20 25 30 Lys Ala Ala Pro Phe Asn Gly Thr Met Met Gln Tyr Phe Glu Trp Tyr Leu Pro Asp Asp Gly Thr Leu Trp Thr Lys Val Ala Asn Glu Ala 50 60 Tyr Lys Gly Thr Ser Arg Ser Asp Val Gly Tyr Gly Val Tyr Asp Leu 85 90 95 Tyr Asp Leu Gly Glu Phe Asn Gln Lys Gly Thr Val Arg Thr Lys Tyr Gly Thr Lys Ala Gln Tyr Leu Gln Ala Ile Gln Ala Ala His Ala Ala 115 120 125 Gly Met Gln Val Tyr Ala Asp Val Val Phe Asp His Lys Gly Gly Ala 130 135 140Asp Gly Thr Glu Trp Val Asp Ala Val Glu Val Asn Pro Ser Asp Arg 145 150 155 160 Asn Gln Glu Ile Ser Gly Thr Tyr Gln Ile Gln Ala Trp Thr Lys Phe 165 170 175 Asp Phe Pro Gly Arg Gly Asn Thr Tyr Ser Ser Phe Lys Trp Arg Trp 180 185 190 Tyr His Phe Asp Gly Val Asp Trp Asp Glu Ser Arg Lys Leu Ser Arg 195 200 205Ile Tyr Lys Phe Arg Gly Ile Gly Lys Ala Trp Asp Trp Glu Val Asp 210 215 220 Thr Glu Asn Gly Asn Tyr Asp Tyr Leu Met Tyr Ala Asp Leu Asp Met Asp His Pro Glu Val Val Thr Glu Leu Lys Asn Trp Gly Lys Trp Tyr 245 250 255 Val Asn Thr Thr Asn Ile Asp Gly Phe Arg Leu Asp Gly Leu Lys His 260 265 270 Ile Lys Phe Ser Phe Phe Pro Asp Trp Leu Ser Tyr Val Arg Ser Gln 275 280 285 Thr Gly Lys Pro Leu Phe Thr Val Gly Glu Tyr Trp Ser Tyr Asp Ile 290 295 300 Asn Lys Leu His Asn Tyr Ile Thr Lys Thr Asn Gly Thr Met Ser Leu 305 310 315 320 Phe Asp Ala Pro Leu His Asn Lys Phe Tyr Thr Ala Ser Lys Ser Gly 325 330 335 Gly Ala Phe Asp Met Arg Thr Leu Met Thr Asn Thr Leu Met Lys Asp 340 345 350 Gln Pro Thr Leu Ala Val Thr Phe Val Asp Asn His Asp Thr Asn Pro

Ala Lys Arg Cys Ser His Gly Arg Pro Trp Phe Lys Pro Leu Ala Tyr 375

Ala Phe Ile Leu Thr Arg Gln Glu Gly Tyr Pro Cys Val Phe Tyr Gly 385

Asp Tyr Tyr Gly Ile Pro Gln Tyr Asn Ile Pro Ser Leu Lys Ser Lys 400

Asp Tyr Tyr Gly Leu Leu Ile Ala Arg Arg Asp Tyr Ala Tyr Gly Thr Gln 425

His Asp Tyr Leu Asp His Ser Asp Ile Ile Gly Trp Thr Arg Glu Gly 435

Val Thr Glu Lys Pro Gly Ser Gly Leu Ala Ala Leu Ile Thr Asp Gly 455

Val Thr Glu Arg Ser Lys Trp Met Tyr Val Gly Lys Gln His Ala Gly Lys 465

Val Phe Tyr Asp Leu Thr Gly Asn Arg Ser Asp Thr Val Thr Ile Asn 485

Ser Asp Gly Trp Gly Glu Phe Lys Val Asp Gly Gly Ser Val Ser Val Ser Val Fro Yal Fro Arg Lys Thr Val Sor Thr Ile Ala Sin Tyr Val Sin Fro Sin Fr

#### (2) INFORMATION FOR SEQ ID NO:36:

- (i) SEQUENCE CHARACTERISTICS:
  - (A) LENGTH: 483 amino acids
  - (B) TYPE: amino acid
  - (C) STRANDEDNESS: single
    (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: protein
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:36:
- Ala Asn Leu Asn Gly Thr Leu Met Gln Tyr Phe Glu Trp Tyr Met Pro
- Asn Asp Gly Cln His Trp Lys Arg Leu Gln Asn Asp Ser Ala Tyr Leu 20 25 30
- Ala Glu His Gly Ile Thr Ala Val Trp Ile Pro Pro Ala Tyr Lys Gly 35 40 45
- Thr Ser Gln Ala Asp Val Gly Tyr Gly Ala Tyr Asp Leu Tyr Asp Leu 50 55
- Gly Glu Phe His Gln Lys Gly Thr Val Arg Thr Lys Tyr Gly Thr Lys 65 70 75 80
- Gly Glu Leu Gln Ser Ala Ile Lys Ser Leu His Ser Arg Asp Ile Asn 85 90 95

Val Tyr Gly Asp Val Val Ile Asn His Lys Gly Gly Ala Asp Ala Thr 100 105 110 Glu Asp Val Thr Ala Val Glu Val Asp Pro Ala Asp Arg Asn Arg Val Ile Ser Gly Glu His Leu Ile Lys Ala Trp Thr His Phe His Phe Pro 130 135 140 Gly Arg Gly Ser Thr Tyr Ser Asp Phe Lys Trp His Trp Tyr His Phe 145 150 155 160 Asp Gly Thr Asp Trp Asp Glu Ser Arg Lys Leu Asn Arg Ile Tyr Lys 165 170 175 Phe Gln Gly Lys Ala Trp Asp Trp Glu Val Ser Asn Glu Asn Gly Asn 180 185 190 Tyr Asp Tyr Leu Thr Tyr Ala Asp Ile Asp Tyr Asp His Pro Asp Val Ala Ala Glu Ile Lys Arg Trp Gly Thr Trp Tyr Ala As<br/>n Glu Leu Gl<br/>n 210 220 Leu Asp Gly Phe Arg Leu Asp Ala Val Lys His Ile Lys Phe Ser Phe 225 230 235 240 Leu Arg Asp Trp Val Asn His Val Arg Glu Lys Thr Gly Lys Glu Met 245 250 255 Phe Thr Val Ala Glu Tyr Trp Gln Asn Asp Leu Gly Ala Leu Glu Asn 260 265 270 Tyr Leu Asn Lys Thr Asn Phe Asn His Ser Val Phe Asp Val Pro Leu 275 280 285 His Tyr Gln Phe His Ala Ala Ser Thr Gln Gly Gly Gly Tyr Asp Met 290 295 300 Arg Lys Leu Leu Asn Gly Thr Val Val Ser Lys His Pro Leu Lys Ser 305 310 315 320 Val Thr Phe Val Asp Asn His Asp Thr Gln Pro Gly Gln Ser Leu Glu 325 330 335 Ser Thr Val Gln Thr Trp Phe Lys Pro Leu Ala Tyr Ala Phe Ile Leu 340 345 350 Thr Arg Glu Ser Gly Tyr Pro Gln Val Phe Tyr Gly Asp Met Tyr Gly 355 360 365 Thr Lys Gly Asp Ser Gln Arg Glu Ile Pro Ala Leu Lys His Lys Ile 370 375 380 Glu Pro Ile Leu Lys Ala Arg Lys Gln Tyr Ala Tyr Gly Ala Gln His 385 390 395 400 Asp Tyr Phe Asp His His Asp Ile Val Gly Trp Thr Arg Glu Gly Asp 405 410 415 Ser Ser Val Ala Asn Ser Gly Leu Ala Ala Leu Ile Thr Asp Gly Pro Gly Gly Ala Lys Arg Met Tyr Val Gly Arg Gln Asn Ala Gly Glu Thr 435 440 445 Trp His Asp Ile Thr Gly Asn Arg Ser Glu Pro Val Val Ile Asn Ser

Glu Gly Trp Gly Glu Phe His Val Asn Gly Gly Ser Val Ser Ile Tyr 465  $\phantom{\bigg|}470\phantom{\bigg|}475\phantom{\bigg|}$ 

Val Gln Arg

- (2) INFORMATION FOR SEQ ID NO:37:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 487 amino acids
      (B) TYPE: amino acid
    - (C) STRANDEDNESS: single
    - (D) TOPOLOGY: linear
  - (ii) MOLECULE TYPE: protein
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:37:

Ala Ala Ala Ala Asn Leu Asn Gly Thr Leu Met Gln Tyr Phe Glu 1 10 15

Trp Tyr Met Pro Asn Asp Gly Gln His Trp Lys Arg Leu Gln Asn Asp 20 25 30

Ser Ala Tyr Leu Ala Glu His Gly Ile Thr Ala Val Trp Ile Pro Pro  $\frac{35}{40}$ 

Ala Tyr Lys Gly Thr Ser Gln Ala Asp Val Gly Tyr Gly Ala Tyr Asp 50 60

Leu Tyr Asp Leu Gly Glu Phe His Gln Lys Gly Thr Val Arg Thr Lys 65 70 80

Tyr Gly Thr Lys Gly Glu Leu Gln Ser Ala Ile Lys Ser Leu His Ser 85 90 95

Arg Asp Ile Asn Val Tyr Gly Asp Val Val Ile Asn His Lys Gly Gly 100 105 110

Ala Asp Ala Thr Glu Asp Val Thr Ala Val Glu Val Asp Pro Ala Asp
115
120
125

Arg Asn Arg Val Ile Ser Gly Glu His Leu Ile Lys Ala Trp Thr His 130 140

Phe His Phe Pro Gly Arg Gly Ser Thr Tyr Ser Asp Phe Lys Trp His 145 155 160

Trp Tyr His Phe Amp Gly Thr Amp Trp Amp Glu Ser Arg Lys Leu Amn 165

Arg Ile Tyr Lys Phe Gln Gly Lys Ala Trp Asp Trp Glu Val Ser Asn 185 190

Glu Asn Gly Asn Tyr Asp Tyr Leu Met Tyr Ala Asp Ile Asp Tyr Asp 195 200 205

His Pro Asp Val Ala Ala Glu Ile Lys Arg Trp Gly Thr Trp Tyr Ala 210 215 220

Asn Glu Leu Gln Leu Asp Gly Phe Arg Leu Asp Ala Val Lys His Ile 225 230 235 240

Lys Phe Ser Phe Leu Arg Asp Trp Val Asn His Val Arg Glu Lys Thr 245 250 255

Gly Lys Glu Met Phe Thr Val Ala Glu Tyr Trp Gln Asn Asp Leu Gly Ala Leu Glu Asn Tyr Leu Asn Lys Thr Asn Phe Asn His Ser Val Phe 275 280 285 Asp Val Pro Leu His Tyr Gln Phe His Ala Ala Ser Thr Gln Gly Gly Gly Tyr Asp Met Arg Lys Leu Leu Asn Gly Thr Val Val Ser Lys His Pro Leu Lys Ser Val Thr Phe Val Asp Asn His Asp Thr Gln Pro Gly Gln Ser Leu Glu Ser Thr Val Gln Thr Trp Phe Lys Pro Leu Ala Tyr 340 345 350 Ala Phe Ile Leu Thr Arg Glu Ser Gly Tyr Pro Gln Val Phe Tyr Gly Asp Met Tyr Gly Thr Lys Gly Asp Ser Gln Arg Glu Ile Pro Ala Leu 370 375 380 Lys His Lys Ile Glu Pro Ile Leu Lys Ala Arg Lys Gln Tyr Ala Tyr 385 390 395 400 Gly Ala Gln His Asp Tyr Phe Asp His His Asp Ile Val Gly Trp Thr 405 410 415 Arg Glu Gly Asp Ser Ser Val Ala Asn Ser Gly Leu Ala Ala Leu Ile 420 425 430 Thr Asp Gly Pro Gly Gly Ala Lys Arg Met Tyr Val Gly Arg Gln Asn 435 440 445Ala Gly Glu Thr Trp His Asp Ile Thr Gly Asn Arg Ser Glu Pro Val 450 460Val Ile Asn Ser Glu Gly Trp Gly Glu Phe His Val Asn Gly Gly Ser Val Ser Ile Tyr Val Gln Arg

- (2) INFORMATION FOR SEQ ID NO:38:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 32 amino acids
    - (B) TYPE: amino acid
    - (C) STRANDEDNESS: single
    - (D) TOPOLOGY: linear
  - (ii) MOLECULE TYPE: protein
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:38:
  - Met Lys Gln Gln Lys Arg Leu Thr Ala Arg Leu Leu Thr Leu Leu Phe  $1 \hspace{1cm} 5 \hspace{1cm} 10 \hspace{1cm} 15$
  - Ala Leu Ile Phe Leu Leu Pro His Ser Ala Ala Ala Ala Ala Asn Leu 20 25 30
- (2) INFORMATION FOR SEQ ID NO:39:

- (i) SEQUENCE CHARACTERISTICS:
  - (A) LENGTH: 33 amino acids
  - (B) TYPE: amino acid
    - (C) STRANDEDNESS: single
    - (D) TOPOLOGY: linear
  - (ii) MOLECULE TYPE: protein
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:39:
  - Met Arg Ser Lys Thr Leu Trp Ile Ser Leu Leu Phe Ala Leu Thr Leu
  - Ile Phe Thr Met Ala Phe Ser Asn Met Ser Ala Gln Ala Ala Gly Lys 20 25 30

Ser

- (2) INFORMATION FOR SEQ ID NO:40:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 35 amino acids
    - (B) TYPE: amino acid (C) STRANDEDNESS: single
    - (D) TOPOLOGY: linear
  - (ii) MOLECULE TYPE: protein
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:40:
  - Met Arg Ser Lys Thr Leu Trp Ile Ser Leu Leu Phe Ala Leu Thr Leu 1 5 10 15
  - Ile Phe Thr Met Ala Phe Ser Asn Met Ser Ala Gln Ala Ala Ala Ala 20 25 30

Ala Ala Asn 35

- (2) INFORMATION FOR SEQ ID NO:41:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 32 amino acids
    - (B) TYPE: amino acid
    - (C) STRANDEDNESS: single (D) TOPOLOGY: linear
  - (ii) MOLECULE TYPE: protein
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:41:
  - Met Arg Ser Lys Thr Leu Trp Ile Ser Leu Leu Phe Ala Leu Thr Leu 1 5 10 15
  - The Phe Thr Met Ala Phe Ser Asn Met Ser Ala Gln Ala Ala Asn Leu  $20 \hspace{1cm} 25 \hspace{1cm} 30 \hspace{1cm}$
- (2) INFORMATION FOR SEQ ID NO:42:
  - (i) SEQUENCE CHARACTERISTICS:
    (A) LENGTH: 33 base pairs

(B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
(ii) MOLECULE TYPE: DNA (genomic)	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:42:	
CACCTAATTA AAGCTTTCAC ACATTTTCAT TTT	3
(2) INFORMATION FOR SEQ ID NO:43:	
(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 33 base pairs (B) TEST and Lake acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
(ii) MOLECULE TYPE: DNA (genomic)	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:43:	
CACCTAATTA AAGCTTACAC ACATTTTCAT TTT	3:
(2) INFORMATION FOR SEQ ID NO:44:	
(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 66 base pairs (B) TTPE: nucleic acid (C) STRANDEDMESS: single (D) TOPCLOOY: linear	
(ii) MOLECULE TYPE: DNA (genomic)	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:44:	
CCGCGTAATT TCCGGAGAAC ACCTAATTAA AGCCGCAACA CATTTCATT TTCCCGGGCG	6
CGGCAG	6
(2) INFORMATION FOR SEQ ID NO:45:	
(i) SEQUENCE CHARACTERISTICS:  (A) LENGTH: 42 base pairs  (B) TYPE: nucleic acid  (C) STRANDEDMESS: single  (D) TOPOLOGY: linear	
(ii) MOLECULE TYPE: DNA (genomic)	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:45:	
CCGGAGAACA CCTAATTAAA GCCCTAACAC ATTTTCATTT TC	4
(2) INFORMATION FOR SEQ ID NO:46:	
(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 42 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single	
57 DIFE 28	
SUBSTITUTE SHEET (RULE 26)	

(D) TOPOLOGY: linear	
(ii) MOLECULE TYPE: DNA (genomic)	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:46:	
CCGGAGAACA CCTAATTAAA GCCCACACAC ATTTTCATTT TC	42
(2) INFORMATION FOR SEQ ID NO:47:	
(i) SEQUENCE CHARACTERISTICS: (A) LENCTH: 42 base pairs (B) TTPS: nucleic acid (C) STRANDEDNESS: single (D) TOPCLOCY: linear	
(ii) MOLECULE TYPE: DNA (genomic)	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:47:	
CCGGAGAACA CCTAATTAAA GCCTGCACAC ATTTTCATTT TC	42
(2) INFORMATION FOR SEQ ID NO:48:	
(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 24 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOFOLOGY: linear	
(ii) MOLECULE TYPE: DNA (genomic)	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:48:	
GATGCAGTAT TTCGAACTGG TATA	24
(2) INFORMATION FOR SEQ ID NO:49:	
(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 26 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPCLOCY: linear	
(ii) MOLECULE TYPE: DNA (genomic)	
(xi) SEQUENCE DESCRIPTION: SEQ ID No:49:	
TGCCCAATGA TGGCCAACAT TGGAAG	26
(2) INFORMATION FOR SEQ ID NO:50:	
(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 24 base pairs (B) TYPE: nucleic acid (C) STRANDEDMESS: single (D) TOPOLOGY: linear	

(ii) MOLECULE TYPE: DNA (genomic)

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:50:	
CGAATGGTAT GCTCCCAATG ACGG	24
(2) INFORMATION FOR SEQ ID NO:51:	
(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 24 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPLOGY: linear	
(ii) MOLECULE TYPE: DNA (genomic)	
•	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:51:	
CGAATGGTAT CGCCCCAATG ACGG	24
(2) INFORMATION FOR SEQ ID NO:52:	
(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 24 base pairs (B) TYPE: nucleic acid (C) STRANDEONESS: single (D) TOPOLOGY: linear	
(ii) MOLECULE TYPE: DNA (genomic)	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:52:	
CGAATGGTAT AATCCCAATG ACGG	24
(2) INFORMATION FOR SEQ ID NO:53:	
(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 24 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
(ii) MOLECULE TYPE: DNA (genomic)	
,	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:53:	
CGAATGGTAT GATCCCAATG ACGG	24
(2) INFORMATION FOR SEQ ID NO:54:	
(i) SEQUENCE CHARACTERISTICS: (A) LENCTH: 24 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
(ii) MOLECULE TYPE: DNA (genomic)	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:54:	

(2) INFORMATION FOR SEQ ID NO:55:	
(i) SEQUENCE CHARACTERISTICS: (a) LENOTH: 24 base pairs (b) TVFs: nucleic acid (c) STRANDEDNESS: single (b) TOPOLOGY: linear	
(ii) MOLECULE TYPE: DNA (genomic)	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:55:	
CGAATGGTAT AAACCCAATG ACGG	24
(2) INFORMATION FOR SEQ ID NO:56:	
(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 24 base pairs (f) LYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
(ii) MOLECULE TYPE: DNA (genomic)	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:56:	
CGAATGGTAT CCGCCCAATG ACGG	24
(2) INFORMATION FOR SEQ ID NO:57:	
(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 24 base pairs (B) TYF: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
(ii) MOLECULE TYPE: DNA (genomic)	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:57:	
CGAATGGTAT TCTCCCAATG ACGG	24
(2) INFORMATION FOR SEQ ID NO:58:	
(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 24 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPCLOGY: linear	
(ii) MOLECULE TYPE: DNA (genomic)	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:58:	
CGAATGGTAC ACTCCCAATG ACGG	24
(2) INFORMATION FOR SEQ ID NO:59:	
(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 24 base pairs	

(B) TYPE: nucleic acid (C) STRANDEDMESS: single (D) TOPOLOGY: linear	
(ii) MOLECULE TYPE: DNA (genomic)	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:59:	
CGAATGGTAT GTTCCCAATG ACGG	24
(2) INFORMATION FOR SEQ ID NO:60:	
(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 24 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
(ii) MOLECULE TYPE: DNA (genomic)	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:60:	
CGAATGGTAT TGTCCCAATG ACGG	24
(2) INFORMATION FOR SEQ ID NO:61:	
(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 24 base pairs (8) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
(ii) MOLECULE TYPE: DNA (genomic)  (xi) SEQUENCE DESCRIPTION: SEQ ID No:61:	
, , -	24
(2) INFORMATION FOR SEQ ID NO:62:	
(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 24 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
(ii) MOLECULE TYPE: DNA (genomic)	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:62:	
CGAATGGTAT GAACCCAATG ACGG	24
(2) INFORMATION FOR SEQ ID No:63:	
(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 24 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	

(ii) MOLECULE TYPE: DNA (genomic)	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:63:	
CGAATGGTAT GGTCCCAATG ACGG	24
(2) INFORMATION FOR SEQ ID NO:64:	
(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 24 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPGLOOF: linear	
(ii) MOLECULE TYPE: DNA (genomic)	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:64:	
CGAATGGTAT ATTCCCAATG ACGG	24
(2) INFORMATION FOR SEQ ID NO:65:	
(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 24 base pairs (B) TYPE: nucleic acid (C) STRANDEDMESS: single (D) TOPOLOGY: linear	
(ii) MOLECULE TYPE: DNA (genomic)	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:65:	
CGAATGGTAT TTTCCCAATG ACGG	24
(2) INFORMATION FOR SEQ ID NO:66:	
(i) SEQUENCE CHARACTERISTICS: (A) LENCTH: 24 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
(ii) MOLECULE TYPE: DNA (genomic)	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:66:	
CGAATGGTAC TGGCCCAATG ACGG	24
(2) INFORMATION FOR SEQ ID NO:67:	
(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 24 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
(ii) MOLECULE TYPE: DNA (genomic)	

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(Xi)	SEQUENCE	DESCRIPTION:	SEQ	ID	NO:67:
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#### CGAATGGTAT TATCCCAATG ACGG

(2) INFORMATION FOR SEQ ID NO:68:

- (i) SEQUENCE CHARACTERISTICS:
  - (A) LENGTH: 24 base pairs
  - (B) TYPE: nucleic acid (C) STRANDEDNESS: single
  - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: DNA (genomic)
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:68:

CCGTCATTGG GACTACGTAC CATT

24

24

#### WHAT IS CLAIMED IS:

A mutant alpha-amylase that is the expression product of a mutated DNA
sequence encoding an alpha-amylase, the mutated DNA sequence being derived from a
precursor alpha-amylase by the deletion or substitution of one or more oxidizable amino
acids selected from the group consisting of methionine, tryptophan, cysteine and
tyrosine, in the precursor alpha-amylase.

- A mutant alpha-amylase of Claim 1 wherein the oxidizable amino acid to be deleted or substituted is a methionine in the precursor alpha-amylase equivalent to +8, +15, +197, +256, +304, +366 or +438 in *Bacillus licheniformis* alpha-amylase.
- A mutant alpha-amylase of Claim 2 wherein the substitution or deletion is at a position equivalent to M+197 in B. licheniformis alpha-amylase.
- 4. A mutant alpha-amylase of Claim 3 wherein an amino acid selected from the group consisting of alanine, isoleucine, threonine and cysteine is substituted for methionine at a position equivalent to +197 in B. licheniformis alpha-amylase.
- The mutant alpha-amylase of Claim 4 which is M197T.
- A mutant alpha-amylase of Claim 2 wherein the substitution or deletion is at a
  position equivalent to M+15 in B. licheniformis alpha-amylase.
- A mutant alpha-amylase of Claim 6 wherein an amino acid selected from the group consisting of leucine, threonine, asparagine, aspartate, serine, valine and isoleucine is substituted for methionine at a position equivalent to +15 in B.

licheniformis alpha-amylase.

The mutant alpha-amylase of Claim 7 which is M15L.

 A mutant alpha-amylase of Claim 1 wherein the oxidizable amino acid to be deleted or substituted is a tryptophan in the precursor alpha-amylase equivalent to any tryptophan in B. licheniformis alpha-amylase as shown in Seq ID No 32.

 A mutant alpha-amylase of Claim 9 wherein the substitution or deletion is at a position equivalent to W138 in B. licheniformis alpha-amylase.

·

 A mutant alpha-amylase of Claim 1 comprising at least two substitutions in a precursor alpha-amylase at positions equivalent to +15, +138 or +197 in B.

licheniformis alpha-amylase.

 A mutant alpha-amylase of Claim 1 wherein the precursor alpha-amylase is a Bacillus alpha-amylase.

13. A mutant alpha-amylase of Claim 12 wherein the precursor is selected from the

group B. licheniformis, B. stearothermophilus, and B. amylolique faciens.

14. A mutant alpha-amylase of Claim 13 wherein the precursor is Bacillus

licheniformis alpha-amylase.

15. A mutant alpha-amylase of Claim 1 wherein the precursor alpha-amylase is a

fungal alpha-amylase.

- 16. DNA encoding the mutant alpha-amylase of Claim 1.
- 17. Expression vectors encoding the DNA of Claim 16.
- 18. Host cells transformed with the expression vector of Claim 17.
- An alpha-amylase comprising an amino acid sequence corresponding to Seq ID
   No 37 or a derivative thereof.
- 20. DNA encoding the alpha-amylase of Claim 19.
- 21. Expression vectors encoding the DNA of Claim 20.
- 22. Host cells transformed with the expression vector of Claim 21.
- 23. A mutant alpha-amylase of Claim 1 having altered oxidative stability comprising a substitution of a different amino acid at a position equivalent to M197 in B. licheniformis alpha-amylase.
- 24. The mutant alpha-amylase of Claim 23 which is M197T.
- 25. A mutant alpha-amylase having enhanced thermal stability, or an enhanced pH performance profile or enhanced oxidative stability, the mutant comprising a substitution of a different amino acid at a position equivalent to M15 in B. licheniformis alpha-amylase.

- 26. The mutant alpha-amylase of Claim 25 which is M15L.
- 27. A detergent composition comprising a mutant alpha-amylase of Claim 1.
- A detergent composition of Claim 27 wherein the mutation is at a position equivalent to M197 in B. licheniformis alpha-amylase.
- A detergent composition of Claim 28 which is a liquid, gel or granular composition.
- A detergent composition of Claim 27 further comprising one or more additional enzyme.
- 31. A starch liquefying composition comprising a mutant alpha-amylase of Claim 1.
- A starch liquefying composition of Claim 31 wherein the mutation is at a position equivalent to M15 in B. licheniformis alpha-amylase.
- 33. A method of liquefying a granular starch slurry from either a wet or dry milling process at a pH of from about 4 to less than about 6 comprising:
  - a) adding an effective amount of an alpha-amylase mutant of Claim 1 to the slurry;
  - b) optionally adding an effective amount of an antioxidant to the slurry; and
  - reacting the slurry for an appropriate time and at an appropriate temperature to liquefy the starch.

34. An improved method of liquefying a granular starch slurry from either a wet or dry milling process at a pH of from about 4 to less than about 6 comprising:

- a) adding an effective amount of an alpha-amylase of Claim 9 to the slurry;
- b) optionally adding an effective amount of an antioxidant to the slurry; and
- reacting the slurry for an appropriate time and at an appropriate temperature to liquefy the starch.

	30 50 CTATTGAATAAATGAGTAGAAAGCGCCATATC			
	90 110 GAAAATGGTACTTGTTAAAAATTCGGAATAT			
	50 170 GGGGAGGAGAATCATGAAACAACAAAAACG M K Q Q K R			
GCTTTACGCCCGATTGCTGACGCTGTTAT	10 230 TTGCGCTCATCTTCTTGCTGCCTCATTCTGC ALIFLPHSA			
250 2 AGCAGCGGCGGCAAATCTTAATGGGACGC	70 290 CTGATGCAGTATTTTGAATGGTACATGCCCAA - M Q Y F E W Y M P N			
	30 350 ACGACTCGGCATATTTGGCTGAACACGGTAT I D S A Y L A E H G I			
TACTGCCGTCTGGATTCCCCCGGCATATAA	90 410 GGGAACGAGCCAAGCGGATGTGGGCTACGG G T S Q A D V G Y G			
TGCTTACGACCTTTATGATTTAGGGGAGTT	.50 470 ITCATCAAAAAGGGACGGTTCGGACAAAGTA HQKGTVRTKY			
CGGCACAAAGGAGAGCTGCAATCTGCG	i10 530 ATCAAAAGTCTTCATTCCCGCGACATTAACGT IKSLHSRDINV			
TTACGGGGATGTGGTCATCAACCACAAAG	570 590 GCGGCGCTGATGCGACCGAAGATGTAACCGC G A D A T E D V T A			
GGTTGAAGTCGATCCCGCTGACCGCAAC	630 CGCGTAATTTCAGGAGAACACCTAATTAAAGC R V I S G E H L I K A			
CTGGACACATTTTCATTTTCCGGGGCGCGCG	890 710 GGCAGCACATACAGCGATTTTAAATGGCATTG GSTYSDFKWHW			
GTACCATTTTGACGGAACCGATTGGGACG	750 770 GAGTCCCGAAAGCTGAACCGCATCTATAAGT1 ESRKLNRIYKF			
TCAAGGAAAGGCTTGGGATTGGGAAGTT	310 830 TCCAATGAAAACGGCAACTATGATTATTTGAT S N E N G N Y D Y L M			
FIG1A				

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850 870 GTATGCCGACATCGATTATGACCATCCTGATGTCGCAGCAGA	890
Y A D I D Y D H P D V A A E	I K R W G T
910 930	950
TTGGTATGCCAATGAACTGCAATTGGACGGTTTCCGTCTTGA	
	AVKHIK
970 990 ATTTTCTTTTTGCGGGATTGGGTTAATCATGTCAGGGAAAAA	1010
F S F L R D W V N H V R E K	T G K E M F
1030 1050	1070
TACGGTAGCTGAATATTGGCAGAATGACTTGGGCGCTCTGGA	
1090 1110	1130
AAATTTTAATCATTCAGTGTTTGACGTGCCGCTTCATTATCAG	
NFNHSVFDVPLHYQ	FHAAST
1150 1170	1190
ACAGGGAGGCGGCTATGATATGAGGAAATTGCTGAACGGTAC Q G G Y D M R K L L N G T	V V S K H P
1210 1230	1250
GTTGAAATCGGTTACATTTGTCGATAACCATGATACACAGCCC	G O S I E S
1270 1290	1310
GACTGTCCAAACATGGTTTAAGCCGCTTGCTTACGCTTTTAT	
TVQTWFKPLAYAFI	L T R E S G
1330 1350	1370
ATACCCTCAGGTTTTCTACGGGGATATGTACGGGACGAAAGG Y P Q V F Y G D M Y G T K G	D S Q R E I
1390 1410	1430
TCCTGCCTTGAAACACAAAATTGAACCGATCTTAAAACGCAG	BAAAACAGTATGCGTACGG
1450 1470	1490
AGCACAGCATGATTATTTCGACCACCATGACATTGTCGGCTGC	GACAAGGGAAGGCGACAG
AQHDYFDHHDIVGW	TREGDS
1510 1530	1550
CTCGGTTGCAAATTCAGGTTTGGCGGCATTAATAACAGACGG, S V A N S G L A A L I T D G	ACCCGGTGGGGCAAAGCG P G G A K R
1570 1590	1610
AATGTATGTCGGCCGGCAAAACGCCGGTGAGACATGGCATG. M Y V G R Q N A G E T W H D	ACATTACCGGAAACCGTTC
1630 1650	1670
GGAGCCGGTTGTCATCAATTCGGAAGGCTGGGGAGAGTTTCA	TODAY A A COCCOCCTOCCT

# FIG.\_1B

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# SUBSTITUTE SHEET (RULE 26)

1810 1830 1850 GTGTCATCAGCCCTCAGGAAGGACTTGCTGACAGTTTGAATCGCATAGGTAAGGCGGGGA

1870 1890 1910 TGAAATGGCAACGTTATCTGATGTAGCAAAGAAAGCAAATGTGTCGAAAATGACGGTATC

1930 1950 GCGGGTGATCAATCATCCTGAGACTGTGACGGATGAATTGAAAAAGCT

FIG.\_1C

FIG.\_1A
FIG.\_1B
FIG.\_1C

50 ANLNGTLMQYFEWYMPNDGOHWKRLONDSAYLAEHGITAVWIPPAYKGTSQADVGYGAYD 90 LYDLGEFHQKGTVRTKYGTKGELQSAIKSLHSRDINVYGDVVINHKGGADATEDVTAVEV 130 150 170 DPADRNRVISGEHLIKAWTHFHFPGRGSTYSDFKWHWYHFDGTDWDESRKLNRIYKFQGK 190 210 230  $AWDWEVSNENGNYDYL\underline{M}YADIDYDHPDVAAEIKRWGTWYANELQLDGFRLDAVKHIKFSF$ 250 270 LRDWVNHVREKTGKEMFTVAEYWQNDLGALENYLNKTNFNHSVFDVPLHYQFHAASTQGG 350 310 330 GYDMRKLLNGTVVSKHPLKSVTFVDNHDTQPGQSLESTVQTWFKPLAYAFILTRESGYPQ 370 390 410 VFYGDMYGTKGDSQREIPALKHKIEPILKARKQYAYGAQHDYFDHHDIVGWTREGDSSVA 430 450 NSGLAALITDGPGGAKRMYVGRQNAGETWHDITGNRSEPVVINSEGWGEFHVNGGSVSIY VQR

FIG. 2

<u> </u>	YFEWYMPNDG YFEWYTPNDG YFEWYLPDDG	79 120 KGTVRTKYGT KGTVRTKYGT KGTVRTKYGT	139 180 SGEHLIKAWT SEEYOIKAWT SGTYQIQAWT	197 240 NENGNYDYLM SENGNYDYLM TENGNYDYLM	257 300 VREKTGKEMF VROATGKEMF VRSQTGKPLE	317 360 LDGTVVSKHP LDGTVVSRHP MTNTLMKDQP
Am-Stearo = $B.stearothermophilus$	AANLNGTLMQ	DLYDLGEFHQ	VDPADRNRVI	GGKAWDWEVS	FSFLRDWVNH	OGGGYDMRKL
	TSAVNGTLMQ	DLYDLGEFQQ	VNPANRNOET	EGKAWDWEVS	FSFLRDWVQA	OGGGYDMRRL
	AAPFNGTMMQ	DLYDLGEFNQ	VNPSDRNOEI	IGKAWDWEVD	FSFFPDWLSY	SGGAFDMRTL
Am-Stearo = B.s	SAAA PITK FCPTGRHAKA	SQADVGYGAY SQSDNGYGPY SRSDVGYGVY	DATEDVTAVE DATEDVTAVE DGTEWVDAVE	KLNRIYKF KISRIFKFRG KLSRIYKFRG	FRLDAVKHIK FRIDAAKHIK FRLDGLKHIK	NFNHSVEDVP LHYOFHAAST SFNQSVEDVP LHFNLQAASS NGTMSLEDAP LHNKFYTASK FIF 3A
Am-Amylo = B.amyloliquefaciens	LFALIFLLPH	VWIPPAYKGT	DVVINHKGGA	FDGTDWDESR	WYANELQLDG	NFNHSVFDVP
	LMCTLLFVSL	VWIPPAYKGL	DVVLNHKAGA	FDGADWDESR	WYANELSDLG	SFNQSVFDVP
	LLAFLLTASL	LSLPPAYKGL	<u>DVVFDH</u> KGGA	FDGVDWDESR	WYVNTTNI <u>DG</u>	NGTMSLFDAP
Am-Amylo = B.a	KRLYARLLTL	AYLAEHGITA	LHSRDINVYG	YSDFKWHWYH	VAAEIKRWGT	GALENYLNKT
	RKRTVSFRLV	EHLSDIGITA	LHSRNVQVYG	YSDFKWHWYH	VVAETKKWGI	GKLENYLNKT
	HRIIRKGWMF	NNLSSLGITA	AHAAGMQVYA	YSSFKWRWYH	VVTELKNWGK	NKLHNYITKT
Am-Lich = B.Licheniformis	1 MRGRGNMIQK VLTF	61 OHWKRLONDS OHWKRLONDA TLWTKVANEA	121 KGELQSAIKS KSELQDAIGS KAQYLQAIQA	181 HFHFPGRGST DFRFPGRGNT KFDFPGRGNT	241 YADIDYDHPD YADVDYDHPD YADLDMDHPE	301 TVAEYWQNDL TVAEYWQNNA TVGEYWSYDI
Am-Lich =	Am-Lich	Am-Lich	Am-Lich	Am-Lich	Am-Lich	Am-Lich
	Am-Amylo	Am-Amylo	Am-Amylo	Am-Amylo	Am-Amylo	Am-Amylo
	Am-Stearo	Am-Stearo	Am-Stearo	Am-Stearo	Am-Stearo	Am-Stearo

**377** 420 480 GTKGDSQREI GTKGTSPKEI GI.....PQYNI ITDGPGGAKR ITDGPGGSKR ITDGAGRSKW YPQVFYGDMY (YPQVFYGDMY (YPCVFYGDYY) SVANSGLAAL SAAKSGLAAL EKPGSGLAAL IVGWTREGDS VIGWTREGDS IIGWTREGVT YAFILTRESG YAFILTRESG YAFILTRQEG TVQTWFKPLA Y TVQTWFKPLA Y HGRPWFKPLA Y AQHDYFDHHD PQHDYIDHPD TQHDYLDHSD DTQPGQSLES DTQPGQSLES DTNPAKR..CS LKARKQYAYG LKARKEYAYG LIARRDYAYG LKSVTFVDNH I EKAVTFVENH I TLAVTFVDNH I PALKHKIEPI PSLKDNIEPI PSLKSKIDPL Am-Amylo Am-Stearo Am-Lich Am-Amylo Am-Stearo Am-Lich

SVWVPRKTTV SIYVOR 483 GEFHVNGGSV GEFHVNDGSV GEFKVNGGSV EPVVINSEGW DTVKIGSDGW DTVTINSDGW TWHDITGNRS TWYDITGNRS VFYDLTGNRS 481 MYVGRQNAGE 1 MYAGLKNAGE 1 MYVKGQHAGK 1 Am-Lich Am-Amylo Am-Stearo

540

STIARPITTR

41 559 WTGEFVRWH EPRLVAWP

> Am-Amylo Am-Stearo

Am-Lich

7,\_3 FIG\_3A

-1G.\_3B

ANLNGTLMQYFEWYMPNDGQHWKRLQNDSAYLAEHGITAVWIPPAYKGTSQADVGYGAYD LYDLGEFHQKGTVRTKYGTKGELQSAIKSLHSRDINVYGDVVINHKGGADATEDVTAVEV DPADRNRVISGEHLIKAWTHFHFPGRGSTYSDFKWHWYHFDGTDWDESRKLNRIYKFQGK  $AWDWEVSNENGNYDYL\underline{T}YADIDYDHPDVAAEIKRWGTWYANELQLDGFRLDAVKHIKFSF$ LRDWVNHVREKTGKEMFTVAEYWQNDLGALENYLNKTNFNHSVFDVPLHYQFHAASTQGG GYDMRKLLNGTVVSKHPLKSVTFVDNHDTQPGQSLESTVQTWFKPLAYAFILTRESGYPQ VFYGDMYGTKGDSQREIPALKHKIEPILKARKQYAYGAQHDYFDHHDIVGWTREGDSSVA NSGLAALITDGPGGAKRMYVGRQNAGETWHDITGNRSEPVVINSEGWGEFHVNGGSVSIY VQR

FIG. 4a

AAAA ANLNGTLMQYFEWYMPNDGQHWKRLQNDSAYLAEHGITAVWIPPAYKGTSQADVGYGAYD LYDLGEFHQKGTVRTKYGTKGELQSAIKSLHSRDINVYGDVVINHKGGADATEDVTAVEV DPADRNRVISGEHLIKAWTHFHFPGRGSTYSDFKWHWYHFDGTDWDESRKLNRIYKFQGK AWDWEVSNENGNYDYLMYADIDYDHPDVAAEIKRWGTWYANELQLDGFRLDAVKHIKFSF LRDWVNHVREKTGKEMFTVAEYWQNDLGALENYLNKTNFNHSVFDVPLHYQFHAASTQGG GYDMRKLLNGTVVSKHPLKSVTFVDNHDTQPGQSLESTVQTWFKPLAYAFILTRESGYPQ VFYGDMYGTKGDSOREIPALKHKIEPILKARKQYAYGAQHDYFDHHDIVGWTREGDSSVA NSGLAALITDGPGGAKRMYVGRQNAGETWHDITGNRSEPVVINSEGWGEFHVNGGSVSIY

. FIG.\_4b

VQR

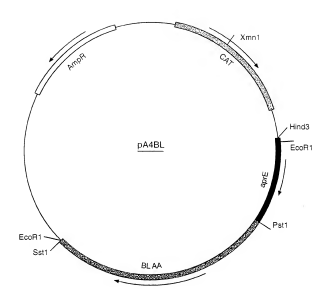


FIG.\_5

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#### SIGNAL SEQUENCE - MATURE PROTEIN JUNCTIONS IN:

B.licheniformis alpha-amylase. (Pstl)

MKQQKRLTARLLTLLFALIFLLPHSAAAAANL.....

B. subtilis alkaline protease aprE.

MRSKTLWISLLFALTLIFTMAFSNMSAQAAGKS.....

B.licheniformis alpha-amylase in pA4BL.

(Pstl)

MRSKTLWISLLFALTLIFTMAFSNMSAQAAAAN.

B.lichenfiormis alpha-amylase in pBLapr.

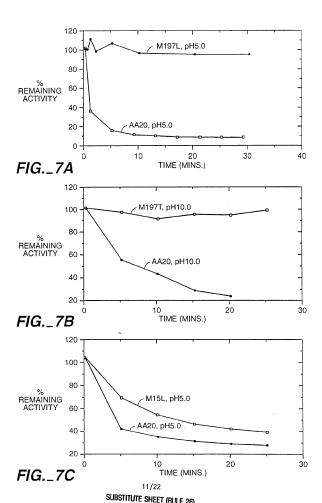
MRSKTLWISLLFALTLIFTMAFSNMSAQAANL.....

(Pstl) indicates the site of the restriction site in the gene.

N-terminus indicates cleavage site between signal peptide and secreted protein.

FIG. 6

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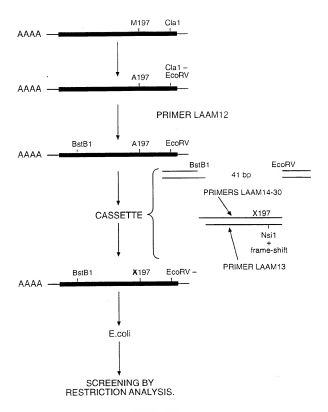
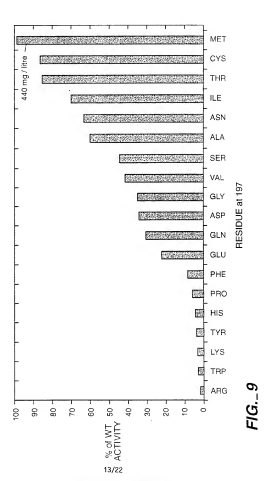


FIG.\_8

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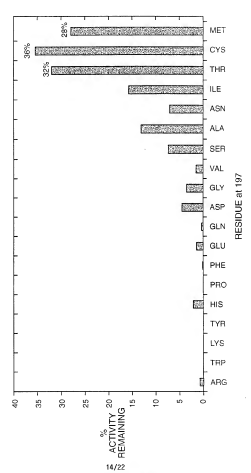


FIG.

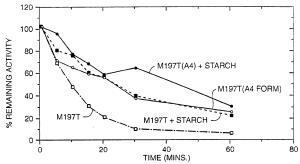


FIG.\_11A

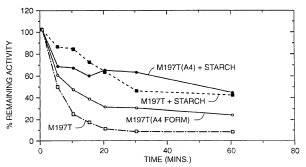


FIG.\_11B

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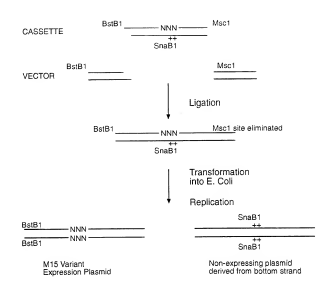
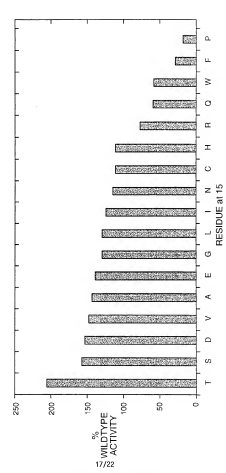
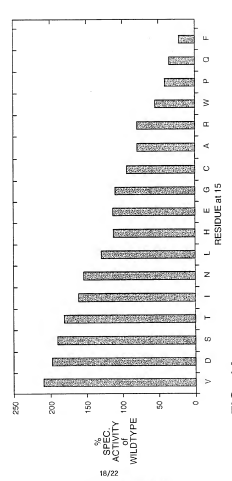


FIG.\_12

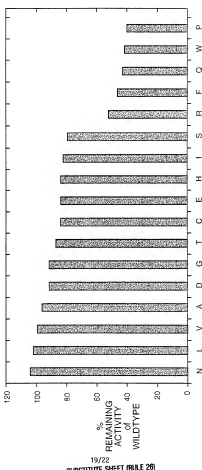
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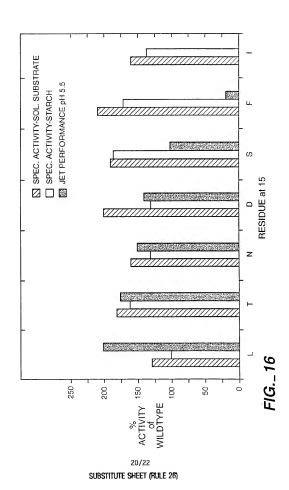
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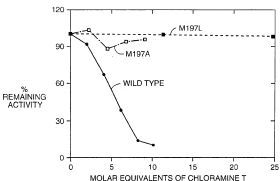
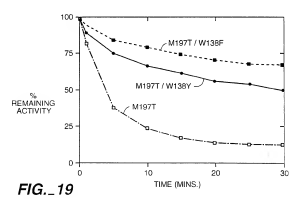


FIG.\_17



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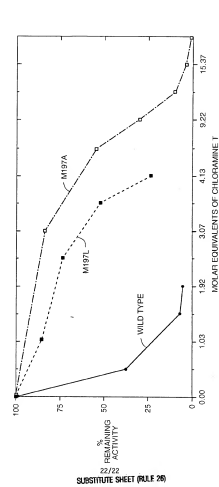


FIG.\_18

## INTERNATIONAL SEARCH REPORT

Intermutional application No. PCT/US 94/01553

A. CLASSIFICATION OF SUBJECT MATTER

IPC: C12N 9/28, C12N 15/56, C11D 3/386
According to International Patent Classification (IPC) or to both national classification and IPC

B. FIELDS SEARCHED

Minimum documentation searched (classification system followed by classification symbols)

IPC : C12N

Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched

Plectronic data base consulted during the international search (name of data base and, where practicable, search terms used)

#### BIOSIS, SCISEARCH

#### C. DOCUMENTS CONSIDERED TO BE RELEVANT

Category*	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.
Y	BIOTECHNOLOGY, Volume 10, 1992, Philippe Joyet et al, "Hyperthermostable variants of a highly thermostable alpha-amylase", page 1579 - page 1583, see fig 4 and page 1582	1-34
Υ	EP, A2, 0410498 (GIST-BROCADES N.V.),	1-34
	30 January 1991 (30.01.91), page 4, line 50 - page 6, claims	
		9
A	FR, A1, 2676456 (INSTITUT NATIONAL DE LA RECHERCHE AGRONOMIQUE), 20 November 1992 (20.11.92), see example 3	1
	<b></b>	
1		1

x Further documents are listed in the continuation of Box C.

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- "E" ertier document but published on or after the international filing date
- document which may throw doubts on priority claim(s) or which is cited to establish the publication date of another citation or other special reason (as specified)
- "O" document referring to an oral disclosure, use, exhibition or other document published prior to the international filing date but later than
- the priority date claimed
- "X" document of particular relevance: the claimed invention cannot be considered novel or cannot be considered to involve an inventive step when the document is taken alone
- document of particular relevance: the claimed invention cannot considered to involve an inventive step when the document is combined with one or more other such documents, such combined.
- being obvious to a person skilled in the art "&" document member of the same patent family

X See patent family annex.

Date of mailing of the international search report

09. 08. 94

Date of the actual completion of the international search

10 June 1994 Varue and mailing address of the international Searching Authorit@uthorized officer

European Patent Office, P.B. 5818 Patentiann 2 NL 2280 HV Rijswik Tel. (+31-70) 340-2040, Tx. 31 651 eoo ni. Fax: (+31-70) 340-3016

CARL-OLOF GUSTAVSSON

## INTERNATIONAL SEARCH REPORT

International application No.
PCT/US 94/01553

1

C (Continuation). DOCUMENTS CONSIDERED TO BE RELEVANT Citation of document, with indication, where appropriate, of the relevant passages Relevant to claim No. Dialog Information Services, File 34, SciSearch Dialog accession no.11267001, Bealinkelly F et al "Studies on the thermostability of the alpha-amylase of bacillus-caldovelox". Applied microbiology and biotechnology, 1991, V36, N3 (DEC), p 332-336 Υ The Journal of Biological Chemistry, Volume 260, No 11, June 1985, David A Estell et al, 1-34 "Engineering an Enzyme by Sitedirected Mutagenesis to Be Resistant to Chemical Oxidation", page 6518 - page 6521, see fig 2 and page 6520 right column Α WO, A1, 9116423 (NOVO NORDISK A/S), 1-34 31 October 1991 (31.10.91), page 2, claims 1-2

Dialog accession no. 11331456, Brosnan MP et al,
"Investigation of the mechanisms of irreversible
thermoinactivation of bacillus-stearothermophilus
alpha-amylase". European journal of biochemistry,
1992, V203, N1-2 (Jan 15), p 225-231

Dialog Information Services, File 34, SciSearch,

X,P WO, A1, 9402597 (NOVO NORDISK A/S), 3 February 1994 (03.02.94), see tables 1-5 and claims

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### SA 8 153

#### INTERNATIONAL SEARCH REPORT Information on patent family members

International application No. 28/05/94 PCT/US 94/01553

Patent document cited in search report		Publication date	Patent family member(s)		Publication date	
EP- <b>A</b> 2-	0410498	30/01/91	AU-B- AU-A- JP-T- WO-A-	638263 5953890 4500756 9100353	24/06/93 17/01/91 13/02/92 10/01/91	
FR-A1-	2676456	20/11/92	NONE			
WO-A1-	9116423	31/10/91	EP-A- US-A-	0528864 5208158	03/03/93 04/05/93	
WO-A1-	9402597	03/02/94	NONE			

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